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(54) METHODS OF TREATING GLOMERULONEPHRITIS ASSOCIATED WITH IGA NEPHROPATHY FOR TACI-IMMUNOGLOBULIN FUSION PROTEINS

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- (60) Division of application No. 13/956,499, filed on Aug. 1, 2013, now Pat. No. 8,815,238, which is a continuation of application No. 13/105,182, filed on May 11, 2011, now Pat. No. 8,524,232, which is a division of application No. 12/605,561, filed on Oct. 26, 2009, now Pat. No. 7,964,711, which is a continuation of application No. 12/359,801, filed on Jan. 26, 2009, now Pat. No. 7,635,767, which is a division of application No. 11/242,294, filed on Oct. 3, 2005, now Pat. No. 7,501,497, which is a continuation of application No. 10/152,363, filed on May 20, 2002, now abandoned.
- (60) Provisional application No. 60/293,343, filed on May 24, 2001.
- (51) **Int. Cl.** A61K 39/00 (2006.01)A61K 45/00 (2006.01)C12P 21/08 (2006.01)C07K 16/24 (2006.01)C07K 14/705 (2006.01)A61K 39/395 (2006.01)A61K 45/06 (2006.01)A61K 38/00 (2006.01)
- (58) Field of Classification Search
 None
 See application file for complete search history.

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(57) ABSTRACT

Molecules that interfere with the binding of a tumor necrosis factor receptor with its ligand, such as a soluble receptor, have proven usefulness in both basic research and as therapeutics. The present invention provides improved soluble transmembrane activator and calcium modulator and cyclophilin ligand-interactor (TACI) receptors.

10 Claims, 7 Drawing Sheets

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30 40 10 20 MSGLGRSRRG GRSRVDQEER FPQGLWTGVA MRSCPEEQYW DPLLGTCMSC 70 80 60 KTICNHOSOR TCAAFCRSLS CRKEQGKFYD HLLRDCISCA SICGOHPKQC 110 120 130 AYFCENKLRS PVNLPPELRR QRSGEVENNS DNSGRYQGLE HRGSEASPAL 170 180 190 PGLKLSADQV ALVYSTLGLC LCAVLCCFLV AVACFLKKRG DPCSCQPRSR 210 220 230 240 250 PRQSPAKSSQ DHAMEAGSPV STSPEPVETC SFCFPECRAP TQESAVTPGT 270 260 280 PDPTCAGRWG CHTRTTVLQP CPHIPDSGLG IVCVPAQEGG PGA

FIGURE 1

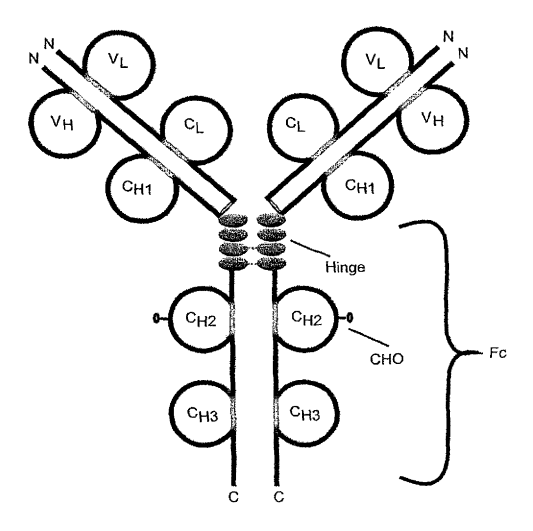


Figure 2

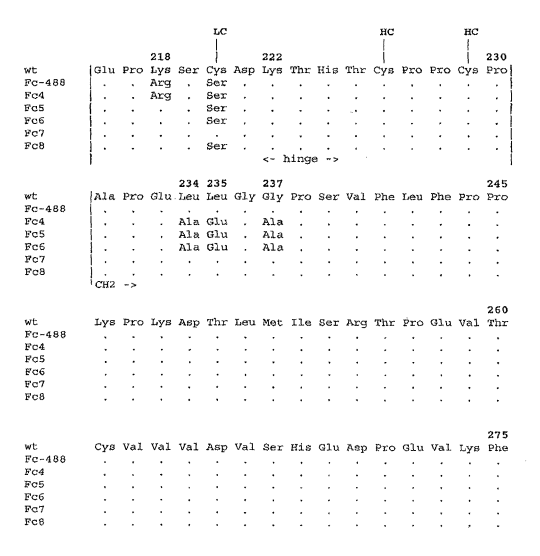


Figure 3A

wt	Asn	Trp	Tvr	Val	Asn	Glv	Val	Glu	Val	His	Asn	Ala	Ĩ.VS	Thr	290 Lvs
Fc-488			-,-			,							1		-3
Fc4	•		•				·	·	•				Ċ	·	
Fc5					·						·	•	·		•
Fc6							Ċ				•	·			
Fc7											•		·		
Fc8															
							297								305
wt	Pro	Arg	Glu	Glu	Gln	ጥነታታ		Ser	Thr	ጥኒታዮ	Δνα	Val	Val	Ser	
Fc-488		ฮ	-	J. U		* y L			* 111	* 7 *					7 CA.L.
Fc4	•		•	·	•	•	•	•	•		•		•	•	
Fc5	·		Ċ			·	·	-	•		·		•	·	
Fc6		·							·					·	·
Fc7		·					Gln	·						·	
Fc8															•
															320
wt	Leu	Thr	Val	Leu	His	Gln	Asp	Trp	Leu	Asn	Gly	Lys	Glu	Tyr	
Fc-488	Leu`	Thr	Val	Leu ,	His	Gln	Asp	Trp	Leu	Asn	Gly	Lys	Glu	Tyr ,	
FC-488 FC4		Thr	Val :	Leu	His :	Gln	Asp :	Trp	Leu :	Asn	Gly	Lys ·	Glu	Tyr	
Fc-488 Fc4 Fc5		Thr	Val	Leu :	His	Gln	Asp :	Trp	Leu	Asn	Gly	Lys	Glu	Tyr	rys
Fc-488 Fc4 Fc5 Fc6		Thr	Val : :	Leu	His	Gln	Asp · ·	Trp	Leu · ·	Asn	Gly : :	Lys · ·	Glu	Tyr	rys
Fc-488 Fc4 Fc5 Fc6 Fc7		Thr	Val	Leu	His	Gln	Asp	Trp	Leu	Asn	Gly	Lys	Glu	Tyr	rys
Fc-488 Fc4 Fc5 Fc6		Thr	Val	Leu	His	Gln	Asp	Trp	Leu	Asn	Gly	Lys	Glu	Туг	· · · rAe
Fc-488 Fc4 Fc5 Fc6 Fc7		Thr	Val.	Leu	His	Gln	Asp	Trp	Leu			Lys	Glu	Туг	Lys ·
Fc-488 Fc4 Fc5 Fc6 Fc7 Fc8		•			•	•				330	331				Lys
Fc-488 Fc4 Fc5 Fc6 Fc7 Fc8		Thr			•	•				330	331				Lys
FC-488 FC4 FC5 FC6 FC7 FC8		•			•	•				330 Ala	331 Pro				Lys
FC-488 FC4 FC5 FC6 FC7 FC8 Wt FC-488 FC4		Lys			•	•				330 Ala	331 Pro				Lys
FC-488 FC4 FC5 FC6 FC7 FC8 Wt FC-488 FC4 FC5		Lys			•	•			Pro	330 Ala Ser	331 Pro			•	Lys
FC-488 FC4 FC5 FC6 FC7 FC8 Wt FC-488 FC4 FC5 FC6		Lys			•	•				330 Ala Ser	331 Pro			•	Lys
FC-488 FC4 FC5 FC6 FC7 FC8 Wt FC-488 FC4 FC5		Lys			•	•			Pro	330 Ala Ser	331 Pro				Lys

Figure 3B

wt Fc-488 Fc4 Fc5 Fc6 Fc7 Fc8	11e	Ser	Lys	Ala	Lys : : CH2		Gln	Pro	Arg	Glu	Pro	Gln	Val	Tyr : : :	350 Thr
						356		358							365
wt	Leu	Pro	Pro	Ser	Arg	Asp	Glu	Leu	Thr	Lys	Asn	Gln	Val	Ser	Leu
Fc-488	•					•	•	•		•	٠				
Fc4	•	•	•	•	•	•	•	•		•	•		•	•	•
Fc5	٠	•	•	•	•	•	٠	•	•	•	•	•	•	•	•
Fc6	•	•	•	•	•	•	•	•	٠	•	•	•	٠	•	٠
Fc7 Fc8	•	•	٠	•	•	•	•	•	•	٠	•	•	٠	٠	•
															380
wt	Thr	Cys	Leu	Val	Lys	Gly	Phe	Tyr	Pro	ser	Asp	Ile	Ala	Val	
Fc-488	Thr	Cys	Leu	Val	rys	Gly	Phe	Tyr	Pro	Ser	Asp •	Ile	Ala	Val	
Fc-488 Fc4	Thr :	Cys	Leu	Val	Lys	Gly :	Phe :	Tyr	Pro	Ser	Asp	Ile	Ala	Val	
Fc-488 Fc4 Fc5	Thr	Cys · ·	Leu	Val	Lys ·	Gly : :	Phe · ·	Tyr	Pro	ser	Asp · ·	Ile :	Ala	Val	
Fc-488 Fc4 Fc5 Fc6	Thr	Cys	Leu	Val	Lys	Gly	Phe · ·	Tyr	Pro	Ser	Asp	Ile	Ala	Val	
FC-488 FC4 FC5 FC6 FC7	Thr	Cys	Leu	Val	Lys	Gly	Phe · ·	Tyr	Pro	ser	Asp	Ile	Ala	Val	
Fc-488 Fc4 Fc5 Fc6	Thr		Leu	Val	Lys	Gly	Phe	Tyr	Pro	ser	Asp · · ·	Ile	Ala	Val	Glu
FC-488 FC4 FC5 FC6 FC7				•	•	•						•	•		Glu
Fc-488 Fc4 Fc5 Fc6 Fc7 Fc8				•	•	•						•	•	Val	Glu
FC-488 FC4 FC5 FC6 FC7 FC8				•	•	•						•	•		Glu
FC-488 FC4 FC5 FC6 FC7 FC8 Wt				•	•	•						•	•		Glu
FC-488 FC4 FC5 FC6 FC7 FC8 Wt FC-488 FC4				•	•	•						•	•		Glu
FC-488 FC4 FC5 FC6 FC7 FC8 Wt FC-488 FC4				•	•	•						•	•		Glu

Figure 3C

															410
wt	Pro	Val	Leu	Asp	Ser	qaA	Gly	Ser	Phe	Phe	Leu	Tyr	Ser	Lys	Leu
Fc-488	•	•	•	•	•	•	•	•	•	•	•	•	•	•	•
Fc4	•	•	•	•	•	-	•	•	•	•	•	٠	•	•	•
Fc5		•	•	•	•	•	•	•		•	•	•	•	•	
Fc6	•	•			•	•	•	•		•	•	•	٠	•	•
Fc7			•	•						٠	•	•	•	•	
Fc8				٠		•		•					•		
															425
wt	Thx	Val	Asp	Lys	Ser	Arg	Trp	Gln	Gln	Gly	Asn	Val	Phe	Ser	Cys
Fc-488		•									•				
Fc4						-									
Fc5			•												
FC6						•			,						-
Fc7									,						
Fc8															
						431	_	1			_				440
wt	ser	Val	Met	нтв	GIU	Ala	ren	His	Asn	His	Тух	Thr	Gin	ŗÀa	Ser
Fc-488	•	•	•	•	٠	•	•	•	٠	•	•	•	•	*	•
Fc4	•	•	•	٠	•	•	•	•	•	•	٠	•	•	•	•
Fc5	•		•	•	•	•	•	•	•	•	٠	٠	٠	•	•
Fc6	•	•	•	•	•	•	•	•	٠	٠	•	•	•	•	•
Fc7	•	•	•	•	•	•	٠	•	•	•	٠	•	•	•	•
Fc8															•
			•	•	•	•	•	•	•	•	•				
			•	•	•	446	•	•	•	•					
wt	Leu	Ser	Leu	Ser	Pro	446 Gly	Lys	***	•	•					
Fc-488	Leu •	Ser	Leu	Ser	Pro		Lys	***	•						
Fc-488 Fc4	Leu :	Ser	Leu	Ser	Pro		Lys	***	•						
Fc-488 Fc4 Fc5	Leu	Ser	Leu	Ser	Pro		•	***	•	·					
Fc-488 Fc4 Fc5 Fc6	Leu	Ser	Leu	Ser	Pro		Lys ***	***	•	·					
Fc-488 Fc4 Fc5	Leu · ·	Ser	Leu	Ser	Pro		•	***	•	•	·				

Figure 3D

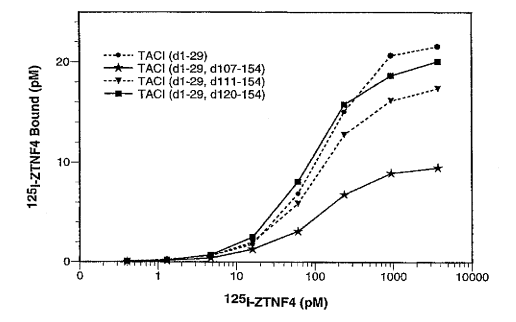


Figure 4

METHODS OF TREATING GLOMERULONEPHRITIS ASSOCIATED WITH IGA NEPHROPATHY FOR TACI-IMMUNOGLOBULIN FUSION **PROTEINS**

CROSS-REFERENCE TO RELATED APPLICATIONS

This application is a divisional of U.S. application Ser. No. 10 13/956,499, filed Aug. 1, 2013, which is a continuation of U.S. application Ser. No. 13/105,182, filed May 11, 2011, now U.S. Pat. No. 8,524,232, which is a divisional of U.S. application Ser. No. 12/605,561, filed Oct. 26, 2009, now U.S. Pat. No. 7,964,711, which is a continuation of U.S. application Ser. No. 12/359,801, filed Jan. 26, 2009, now U.S. Pat. No. 7,635,767, which is a divisional of U.S. application Ser. No. 11/242,294, filed Oct. 3, 2005, now U.S. Pat. No. 7,501,497, which is a continuation of U.S. application Ser. No. 10/152,363, now abandoned, filed May 20, 2002, which claims the benefit of U.S. Provisional Application Ser. No. 60/293,343, filed May 24, 2001, each of which are herein incorporated by reference.

TECHNICAL FIELD

The present invention relates generally to improved fusion proteins comprising a tumor necrosis factor receptor moiety and an immunoglobulin moiety. In particular, the present invention relates to improved TACI-immunoglobulin fusion 30 proteins.

REFERENCE TO A SEQUENCE LISTING SUBMITTED AS A TEXT FILE VIA EFS-WEB

The official copy of the sequence listing is submitted concurrently with the specification as a text file via EFS-Web, in compliance with the American Standard Code for Information Interchange (ASCII), with a file name of 448288seqlist-.txt, a creation date of Jul. 15, 2014, and a size of 100 Kb. The 40 sequence listing filed via EFS-Web is part of the specification and is hereby incorporated in its entirety by reference herein.

BACKGROUND OF THE INVENTION

Cytokines are soluble, small proteins that mediate a variety of biological effects, including the regulation of the growth and differentiation of many cell types (see, for example, Arai et al., Annu. Rev. Biochem. 59:783 (1990); Mosmann, Curr. Opin. Immunol. 3:311 (1991); Paul and Seder, Cell 76:241 50 (1994)). Proteins that constitute the cytokine group include interleukins, interferons, colony stimulating factors, tumor necrosis factors, and other regulatory molecules. For example, human interleukin-17 is a cytokine which stimulates the expression of interleukin-6, intracellular adhesion 55 molecule 1, interleukin-8, granulocyte macrophage colonystimulating factor, and prostaglandin E2 expression, and plays a role in the preferential maturation of CD34+ hematopoietic precursors into neutrophils (Yao et al., J. Immunol. 155:5483 (1995); Fossiez et al., J. Exp. Med. 183:2593 60

Receptors that bind cytokines are typically composed of one or more integral membrane proteins that bind the cytokine with high affinity and transduce this binding event to the cell through the cytoplasmic portions of the certain receptor 65 locations of the cysteine-rich pseudo-repeats are indicated by subunits. Cytokine receptors have been grouped into several classes on the basis of similarities in their extracellular ligand

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binding domains. For example, the receptor chains responsible for binding and/or transducing the effect of interferons are members of the type II cytokine receptor family, based upon a characteristic 200 residue extracellular domain.

Cellular interactions, which occur during an immune response, are regulated by members of several families of cell surface receptors, including the tumor necrosis factor receptor (TNFR) family. The TNFR family consists of a number of integral membrane glycoprotein receptors many of which, in conjunction with their respective ligands, regulate interactions between different hematopoietic cell lineages (see, for example, Cosman, Stem Cells 12:440 (1994); Wajant et al., Cytokine Growth Factor Rev. 10:15 (1999); Yeh et al., Immunol. Rev. 169:283 (1999); Idriss and Naismith, Microsc. Res. Tech. 50:184 (2000)).

One such receptor is TACI, transmembrane activator and CAML-interactor (von Bülow and Bram, Science 228:138 (1997); Bram and von Bülow, U.S. Pat. No. 5,969,102 (1999)). TACI is a membrane bound receptor, which has an extracellular domain containing two cysteine-rich pseudorepeats, a transmembrane domain and a cytoplasmic domain that interacts with CAML (calcium-modulator and cyclophilin ligand), an integral membrane protein located at intracellular vesicles which is a co-inducer of NF-AT activation when overexpressed in Jurkat cells. TACI is associated with B cells and a subset of T cells. Nucleotide sequences that encode TACI and its corresponding amino acid sequence are provided herein as SEQ ID NOs: 1 and 2, respectively

The TACI receptor binds two members of the tumor necrosis factor (TNF) ligand family. One ligand is variously designated as ZTNF4, "BAFF," "neutrokine-α," "BLyS," "TALL-1," and "THANK" (Yu et al., international publication No. WO98/18921 (1998), Moore et al., Science 285:269 (1999); Mukhopadhyay et al., J. Biol. Chem. 274:15978 (1999); Schneider et al., J. Exp. Med. 189:1747 (1999); Shu et al., J. Leukoc. Biol. 65:680 (1999)). The amino acid sequence of ZTNF4 is provided as SEQ ID NO:3. The other ligand has been designated as "ZTNF2," "APRIL" and "TNRF death ligand-1" (Hahne et al., J. Exp. Med. 188:1185 (1998); Kelly et al., Cancer Res. 60:1021 (2000)). The amino acid sequence of ZTNF2 is provided as SEQ ID NO:4. Both ligands are also bound by the B-cell maturation receptor (BCMA) (Gross et al., Nature 404:995 (2000)). The nucleotide and amino acid sequence of BCMA are provided as SEQ ID NO:26 and SEQ ID NO:27, respectively.

The demonstrated in vivo activities of tumor necrosis factor receptors illustrate the clinical potential of soluble forms of the receptor. Soluble forms of the TACI receptor have been generated as immunoglobulin fusion proteins. Initial versions resulted in low-expressing, heterogeneous protein. The heterogeneity was observed at the TACI amino terminus, at the Fc carboxyl terminus, and in the TACI stalk region. A need therefore exists for pharmaceutically useful TACI receptor compositions.

BRIEF SUMMARY OF THE INVENTION

The present invention provides improved TACI-immunoglobulin fusion proteins suitable as therapeutic compounds.

BRIEF DESCRIPTION OF THE DRAWINGS

FIG. 1 shows the amino acid sequence of human TACI. The shading, the transmembrane domain is boxed, and the stalk region is indicated by hash marks.

FIG. **2** is a schematic diagram of an immunoglobulin of the IgG1 subclass. C_L : light chain constant region; C_{H1} , C_{H2} , C_{H3} : heavy chain constant region; V_L : light chain variable region; V_{H^2} : heavy chain variable region; CHO: carbohydrate; N: amino terminus; C: carboxyl terminus.

FIGS. 3A, 3B, 3C, and 3D show a comparison of the wild-type human $\gamma 1$ constant region Fc amino acid sequence with variants Fc-488, Fc4, Fc5, Fc6, Fc7, and Fc8. The C_{H1} domain of the human $\gamma 1$ constant region is not part of the Fc and is therefore not shown. The location of the hinge region, 10 the C_{H2} , and the C_{H3} domains are indicated. The Cys residues normally involved in disulfide bonding to the light chain constant region (LC) and heavy chain constant region (HC) are indicated. A "." symbol indicates identity to wild-type at that position, while "***" indicates the location of the carboxyl terminus, and illustrates the difference in the carboxyl terminus of Fc6 relative to the other Fc versions. Amino acid locations are indicated by EU index positions.

FIG. 4 shows the specific binding of ¹²⁵I-ZTNF4 with various TACI-Fc constructs. The TACI-Fc fusion proteins ²⁰ had TACI moieties that lacked the first 29 amino acid residues of the amino acid sequence of SEQ ID NO:2. One of the fusion proteins had a TACI moiety with an intact stalk region (TACI (d1-29)-Fc5), whereas three of the TACI-Fc fusion proteins had TACI moieties with various deletions in the stalk ²⁵ region (TACI (d1-29, d107-154)-Fc5; TACI (d1-29, d111-154)-Fc5; TACI (d1-29, d120-154)-Fc5). Experimental details are described in Example 4.

DETAILED DESCRIPTION OF THE INVENTION

1. Overview

As described below, the present invention provides transmembrane activator and calcium modulator and cyclophilin 35 ligand-interactor (TACI)-immunoglobulin fusion proteins, and methods for using TACI-immunoglobulin fusion proteins. For example, the present invention provides methods for inhibiting the proliferation of tumor cells, comprising administering to the tumor cells a composition that comprises 40 a TACI-immunoglobulin fusion protein. Such a composition can be administered to cells cultured in vitro. Alternatively, the composition can be a pharmaceutical composition that comprises a pharmaceutically acceptable carrier and a TACIimmunoglobulin fusion protein, and the pharmaceutical com- 45 position can be administered to a subject, which has a tumor. The subject may be a mammalian subject. Administration of the pharmaceutical composition can inhibit, for example, the proliferation of B lymphocytes in a mammalian subject.

The present invention also provides methods for inhibiting 50 ZTNF4 activity in a mammal, comprising administering to the mammal a composition that comprises a TACI-immunoglobulin. The ZTNF4 activity can be associated with various diseases and disorders. For example, a pharmaceutical composition that comprises a TACI-immunoglobulin fusion pro- 55 tein can be used to treat an autoimmune disease, such as systemic lupus erythematosus, myasthenia gravis, multiple sclerosis, insulin dependent diabetes mellitus, Crohn's disease, rheumatoid arthritis, polyarticular-course juvenile rheumatoid arthritis, and psoriatic arthritis. Alternatively, a phar- 60 maceutical composition that comprises a TACIimmunoglobulin can be used to treat a disorder such as asthma, bronchitis, emphysema, and end stage renal failure. A pharmaceutical composition comprising a TACI-immunoglobulin can also be used to treat renal disease, such as glom- 65 erulonephritis, vasculitis, nephritis, amyloidosis, and pyelonephritis, or a disorder, such as neoplasm, chronic

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lymphocytic leukemia, multiple myeloma, non-Hodgkin's lymphoma, post-transplantation lymphoproliferative disease, and light chain gammopathy. In certain cases, the ZTNF4 activity can be associated with T cells. A pharmaceutical composition that comprises a TACI-immunoglobulin can also be used to treat a disease or disorder associated with immunosuppression, graft rejection, graft versus host disease, and inflammation. For example, a pharmaceutical composition that comprises a TACI-immunoglobulin can be used to decrease inflammation, and to treat disorders such as joint pain, swelling, anemia, and septic shock.

The present invention also provides methods for reducing circulating blood levels of ZTNF4 in a mammalian subject, comprising administering to the mammalian subject a pharmaceutical composition that comprises a pharmaceutically acceptable carrier and a TACI-immunoglobulin fusion protein, wherein administration of the pharmaceutical composition reduces the circulating level of ZTNF4 in the blood of the mammalian subject. As an illustration, the administration of such a pharmaceutical composition can reduce circulating blood levels of ZTNF4 by at least 10%, by at least 20%, by at least 10 to 60%, by at least 20 to 50%, or by at least 30 to 40%, compared with the blood level of ZTNF4 prior to the administration of the pharmaceutical composition. Those of skill in the art can measure circulating levels of ZTNF4. Illustrative methods are described in Example 4 and Example 5.

As described below, illustrative TACI-immunoglobulin fusion proteins comprise:

- (a) a TACI receptor moiety that consists of a fragment of a polypeptide that has the amino acid sequence of amino acid residues 30 to 154 of SEQ ID NO:2, wherein the TACI receptor moiety comprises at least one of (i) amino acid residues 34 to 66 of SEQ ID NO:2, and (ii) amino acid residues 71 to 104 of SEQ ID NO:2, and wherein the TACI receptor moiety binds at least one of ZTNF2 or ZTNF4, and
- (b) an immunoglobulin moiety comprising a constant region of an immunoglobulin.

Suitable TACI receptor moieties include: polypeptides that comprise amino acid residues 34 to 66 of SEQ ID NO:2, and amino acid residues 71 to 104 of SEQ ID NO:2; polypeptides that comprise amino acid residues 34 to 104 of SEQ ID NO:2; polypeptides that comprise the amino acid sequence of amino acid residues 30 to 110 of SEQ ID NO:2; and polypeptides that have an amino acid sequence consisting of amino acid residues 30 to 110 of SEQ ID NO:2.

The immunoglobulin moiety of a TACI-immunoglobulin fusion protein can comprise a heavy chain constant region, such as a human heavy chain constant region. An IgG1 heavy chain constant region is one example of a suitable heavy chain constant region. An illustrative IgG1 heavy chain constant region is an IgG1 Fc fragment that comprises C_{H2} , and C_{H3} domains. The IgG1 Fc fragment can be a wild-type IgG1 Fc fragment or a mutated IgG1 Fc fragment, such as the Fc fragment comprising the amino acid sequence of SEQ ID NO:33. One exemplary TACI-immunoglobulin fusion protein is a protein that has an amino acid sequence comprising the amino acid sequence of SEQ ID NO:54.

The TACI-immunoglobulin fusion proteins described herein can be multimers, such as dimers.

The present invention also provides nucleic acid molecules that encode a TACI-immunoglobulin fusion protein. An illustrative nucleotide sequence that encodes a TACI-immunoglobulin fusion protein is provided by SEQ ID NO:53.

The present invention also includes TACI soluble receptors that consist of a fragment of a polypeptide that has the amino acid sequence of amino acid residues 30 to 154 of SEQ ID

NO:2, wherein the TACI soluble receptor comprises at least one of (i) amino acid residues 34 to 66 of SEQ ID NO:2, and (ii) amino acid residues 71 to 104 of SEQ ID NO:2, and wherein the TACI soluble receptor binds at least one of ZTNF2 or ZTNF4. Additional TACI soluble receptors are described herein as suitable TACI receptor moieties for TACI-immunoglobulin fusion proteins. Moreover, TACI soluble receptors can be used in methods described for TACI-immunoglobulin fusion proteins.

These and other aspects of the invention will become evident upon reference to the following detailed description and drawings. In addition, various references are identified below and are incorporated by reference in their entirety.

2. Definitions

In the description that follows, a number of terms are used extensively. The following definitions are provided to facilitate understanding of the invention.

As used herein, "nucleic acid" or "nucleic acid molecule" 20 refers to polynucleotides, such as deoxyribonucleic acid (DNA) or ribonucleic acid (RNA), oligonucleotides, fragments generated by the polymerase chain reaction (PCR), and fragments generated by any of ligation, scission, endonuclease action, and exonuclease action. Nucleic acid mol- 25 ecules can be composed of monomers that are naturallyoccurring nucleotides (such as DNA and RNA), or analogs of naturally-occurring nucleotides (e.g., α -enantiomeric forms of naturally-occurring nucleotides), or a combination of both. Modified nucleotides can have alterations in sugar moieties 30 and/or in pyrimidine or purine base moieties. Sugar modifications include, for example, replacement of one or more hydroxyl groups with halogens, alkyl groups, amines, and azido groups, or sugars can be functionalized as ethers or esters. Moreover, the entire sugar moiety can be replaced with 35 sterically and electronically similar structures, such as azasugars and carbocyclic sugar analogs. Examples of modifications in a base moiety include alkylated purines and pyrimidines, acylated purines or pyrimidines, or other well-known heterocyclic substitutes. Nucleic acid monomers can be 40 linked by phosphodiester bonds or analogs of such linkages. Analogs of phosphodiester linkages include phosphorothioate, phosphorodithioate, phosphoroselenoate, phosphorodiselenoate, phosphoroanilothioate, phosphoranilidate, phosphoramidate, and the like. The term "nucleic acid molecule" 45 also includes so-called "peptide nucleic acids," which comprise naturally-occurring or modified nucleic acid bases attached to a polyamide backbone. Nucleic acids can be either single stranded or double stranded.

The term "complement of a nucleic acid molecule" refers 50 to a nucleic acid molecule having a complementary nucleotide sequence and reverse orientation as compared to a reference nucleotide sequence. For example, the sequence 5' ATGCACGGG 3' (SEQ ID NO:57) is complementary to 5' CCCGTGCAT 3' (SEQ ID NO:58).

The term "contig" denotes a nucleic acid molecule that has a contiguous stretch of identical or complementary sequence to another nucleic acid molecule. Contiguous sequences are said to "overlap" a given stretch of a nucleic acid molecule either in their entirety or along a partial stretch of the nucleic 60 acid molecule.

The term "degenerate nucleotide sequence" denotes a sequence of nucleotides that includes one or more degenerate codons as compared to a reference nucleic acid molecule that encodes a polypeptide. Degenerate codons contain different 65 triplets of nucleotides, but encode the same amino acid residue (i.e., GAU and GAC triplets each encode Asp).

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The term "structural gene" refers to a nucleic acid molecule that is transcribed into messenger RNA (mRNA), which is then translated into a sequence of amino acids characteristic of a specific polypeptide.

An "isolated nucleic acid molecule" is a nucleic acid molecule that is not integrated in the genomic DNA of an organism. For example, a DNA molecule that encodes a growth factor that has been separated from the genomic DNA of a cell is an isolated DNA molecule. Another example of an isolated nucleic acid molecule is a chemically-synthesized nucleic acid molecule that is not integrated in the genome of an organism. A nucleic acid molecule that has been isolated from a particular species is smaller than the complete DNA molecule of a chromosome from that species.

A "nucleic acid molecule construct" is a nucleic acid molecule, either single- or double-stranded, that has been modified through human intervention to contain segments of nucleic acid combined and juxtaposed in an arrangement not existing in nature.

"Linear DNA" denotes non-circular DNA molecules having free 5' and 3' ends. Linear DNA can be prepared from closed circular DNA molecules, such as plasmids, by enzymatic digestion or physical disruption.

"Complementary DNA (cDNA)" is a single-stranded DNA molecule that is formed from an mRNA template by the enzyme reverse transcriptase. Typically, a primer complementary to portions of mRNA is employed for the initiation of reverse transcription. Those skilled in the art also use the term "cDNA" to refer to a double-stranded DNA molecule consisting of such a single-stranded DNA molecule and its complementary DNA strand. The term "cDNA" also refers to a clone of a cDNA molecule synthesized from an RNA template.

A "promoter" is a nucleotide sequence that directs the transcription of a structural gene. Typically, a promoter is located in the 5' non-coding region of a gene, proximal to the transcriptional start site of a structural gene. Sequence elements within promoters that function in the initiation of transcription are often characterized by consensus nucleotide sequences. These promoter elements include RNA polymerase binding sites, TATA sequences, CAAT sequences, differentiation-specific elements (DSEs; McGehee et al., Mol. Endocrinol. 7:551 (1993)), cyclic AMP response elements (CREs), serum response elements (SREs; Treisman, Seminars in Cancer Biol. 1:47 (1990)), glucocorticoid response elements (GREs), and binding sites for other transcription factors, such as CRE/ATF (O'Reilly et al., J. Biol. Chem. 267:19938 (1992)), AP2 (Ye et al., J. Biol. Chem. 269:25728 (1994)), SP1, cAMP response element binding protein (CREB; Loeken, Gene Expr. 3:253 (1993)) and octamer factors (see, in general, Watson et al., eds., Molecular Biology of the Gene, 4th ed. (The Benjamin/Cummings Publishing Company, Inc. 1987), and Lemaigre and Rousseau, Biochem. J. 303:1 (1994)). If a promoter is an inducible 55 promoter, then the rate of transcription increases in response to an inducing agent. In contrast, the rate of transcription is not regulated by an inducing agent if the promoter is a constitutive promoter. Repressible promoters are also known.

A "core promoter" contains essential nucleotide sequences for promoter function, including the TATA box and start of transcription. By this definition, a core promoter may or may not have detectable activity in the absence of specific sequences that may enhance the activity or confer tissue specific activity.

A "regulatory element" is a nucleotide sequence that modulates the activity of a core promoter. For example, a regulatory element may contain a nucleotide sequence that

binds with cellular factors enabling transcription exclusively or preferentially in particular cells, tissues, or organelles. These types of regulatory elements are normally associated with genes that are expressed in a "cell-specific," "tissuespecific," or "organelle-specific" manner.

An "enhancer" is a type of regulatory element that can increase the efficiency of transcription, regardless of the distance or orientation of the enhancer relative to the start site of transcription.

"Heterologous DNA" refers to a DNA molecule, or a population of DNA molecules, that does not exist naturally within a given host cell. DNA molecules heterologous to a particular host cell may contain DNA derived from the host cell species (i.e., endogenous DNA) so long as that host DNA is combined with non-host DNA (i.e., exogenous DNA). For example, a DNA molecule containing a non-host DNA segment encoding a polypeptide operably linked to a host DNA segment comprising a transcription promoter is considered to be a heterologous DNA molecule. Conversely, a heterologous 20 DNA molecule can comprise an endogenous gene operably linked with an exogenous promoter. As another illustration, a DNA molecule comprising a gene derived from a wild-type cell is considered to be heterologous DNA if that DNA molecule is introduced into a mutant cell that lacks the wild-type 25 gene.

A "polypeptide" is a polymer of amino acid residues joined by peptide bonds, whether produced naturally or synthetically. Polypeptides of less than about 10 amino acid residues are commonly referred to as "peptides."

A "protein" is a macromolecule comprising one or more polypeptide chains. A protein may also comprise non-peptidic components, such as carbohydrate groups. Carbohydrates and other non-peptidic substituents may be added to a protein by the cell in which the protein is produced, and will vary with the type of cell. Proteins are defined herein in terms of their amino acid backbone structures; substituents such as carbohydrate groups are generally not specified, but may be present nonetheless.

A peptide or polypeptide encoded by a non-host DNA molecule is a "heterologous" peptide or polypeptide.

An "integrated genetic element" is a segment of DNA that has been incorporated into a chromosome of a host cell after that element is introduced into the cell through human 45 manipulation. Within the present invention, integrated genetic elements are most commonly derived from linearized plasmids that are introduced into the cells by electroporation or other techniques. Integrated genetic elements are passed from the original host cell to its progeny.

A "cloning vector" is a nucleic acid molecule, such as a plasmid, cosmid, or bacteriophage, which has the capability of replicating autonomously in a host cell. Cloning vectors typically contain one or a small number of restriction endonuclease recognition sites that allow insertion of a nucleic 55 acid molecule in a determinable fashion without loss of an essential biological function of the vector, as well as nucleotide sequences encoding a marker gene that is suitable for use in the identification and selection of cells transformed with the cloning vector. Marker genes typically include genes 60 that provide tetracycline resistance or ampicillin resistance.

An "expression vector" is a nucleic acid molecule encoding a gene that is expressed in a host cell. Typically, an expression vector comprises a transcription promoter, a gene, and a transcription terminator. Gene expression is usually 65 placed under the control of a promoter, and such a gene is said to be "operably linked to" the promoter. Similarly, a regula-

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tory element and a core promoter are operably linked if the regulatory element modulates the activity of the core promoter

A "recombinant host" is a cell that contains a heterologous nucleic acid molecule, such as a cloning vector or expression vector. In the present context, an example of a recombinant host is a cell that produces a TACI-Fc fusion protein from an expression vector.

"Integrative transformants" are recombinant host cells, in which heterologous DNA has become integrated into the genomic DNA of the cells.

A "fusion protein" is a hybrid protein expressed by a nucleic acid molecule comprising nucleotide sequences of at least two genes. For example, a TACI-immunoglobulin fusion protein comprises a TACI receptor moiety and an immunoglobulin moiety. As used herein, a "TACI receptor moiety" is a portion of the extracellular domain of the TACT receptor that binds at least one of ZTNF2 or ZTNF4. The phrase an "immunoglobulin moiety" refers to a polypeptide that comprises a constant region of an immunoglobulin. For example, the immunoglobulin moiety can comprise a heavy chain constant region. The term "TACI-Fc" fusion protein refers to a TACI-immunoglobulin fusion protein in which the immunoglobulin moiety comprises immunoglobulin heavy chain constant regions, C_{H2} and C_{H3} .

The term "receptor" denotes a cell-associated protein that binds to a bioactive molecule termed a "ligand." This interaction mediates the effect of the ligand on the cell. In the context of TACI receptor binding, the phrase "specifically binds" or "specific binding" refers to the ability of the ligand to competitively bind with the receptor. For example, ZTNF4 specifically binds with the TACI receptor, and this can be shown by observing competition for the TACI receptor between detectably labeled ZTNF4 and unlabeled ZTNF4.

Receptors can be membrane bound, cytosolic or nuclear; monomeric (e.g., thyroid stimulating hormone receptor, beta-adrenergic receptor) or multimeric (e.g., PDGF receptor, growth hormone receptor, IL-3 receptor, GM-CSF receptor, G-CSF receptor, erythropoietin receptor and IL-6 receptor).
 Membrane-bound receptors are characterized by a multi-domain structure comprising an extracellular ligand-binding domain and an intracellular effector domain that is typically involved in signal transduction. In certain membrane-bound receptors, the extracellular ligand-binding domain and the
 intracellular effector domain are located in separate polypeptides that comprise the complete functional receptor.

In general, the binding of ligand to receptor results in a conformational change in the receptor that causes an interaction between the effector domain and other molecule(s) in the cell, which in turn leads to an alteration in the metabolism of the cell. Metabolic events that are often linked to receptorligand interactions include gene transcription, phosphorylation, dephosphorylation, increases in cyclic AMP production, mobilization of cellular calcium, mobilization of membrane lipids, cell adhesion, hydrolysis of inositol lipids and hydrolysis of phospholipids.

The term "secretory signal sequence" denotes a DNA sequence that encodes a peptide (a "secretory peptide") that, as a component of a larger polypeptide, directs the larger polypeptide through a secretory pathway of a cell in which it is synthesized. The larger polypeptide is commonly cleaved to remove the secretory peptide during transit through the secretory pathway.

An "isolated polypeptide" is a polypeptide that is essentially free from contaminating cellular components, such as carbohydrate, lipid, or other proteinaceous impurities associated with the polypeptide in nature. Typically, a preparation

of isolated polypeptide contains the polypeptide in a highly purified form, i.e., at least about 80% pure, at least about 90% pure, at least about 95% pure, greater than 95% pure, or greater than 99% pure. One way to show that a particular protein preparation contains an isolated polypeptide is by the appearance of a single band following sodium dodecyl sulfate (SDS)-polyacrylamide gel electrophoresis of the protein preparation and Coomassie Brilliant Blue staining of the gel. However, the term "isolated" does not exclude the presence of the same polypeptide in alternative physical forms, such as dimers or alternatively glycosylated or derivatized forms.

The terms "amino-terminal" and "carboxyl-terminal" are used herein to denote positions within polypeptides. Where the context allows, these terms are used with reference to a particular sequence or portion of a polypeptide to denote proximity or relative position. For example, a certain sequence positioned carboxyl-terminal to a reference sequence within a polypeptide is located proximal to the carboxyl terminus of the reference sequence, but is not nec- 20 essarily at the carboxyl terminus of the complete polypeptide.

The term "expression" refers to the biosynthesis of a gene product. For example, in the case of a structural gene, expression involves transcription of the structural gene into mRNA and the translation of mRNA into one or more polypeptides. 25

The term "splice variant" is used herein to denote alternative forms of RNA transcribed from a gene. Splice variation arises naturally through use of alternative splicing sites within a transcribed RNA molecule, or less commonly between separately transcribed RNA molecules, and may result in 30 several mRNAs transcribed from the same gene. Splice variants may encode polypeptides having altered amino acid sequence. The term splice variant is also used herein to denote a polypeptide encoded by a splice variant of an mRNA transcribed from a gene.

As used herein, the term "immunomodulator" includes cytokines, stem cell growth factors, lymphotoxins, co-stimulatory molecules, hematopoietic factors, and synthetic analogs of these molecules.

The term "complement/anti-complement pair" denotes 40 non-identical moieties that form a non-covalently associated, stable pair under appropriate conditions. For instance, biotin and avidin (or streptavidin) are prototypical members of a complement/anti-complement pair. Other exemplary pairs, antibody/antigen (or hapten or epitope) pairs, sense/ antisense polynucleotide pairs, and the like. Where subsequent dissociation of the complement/anti-complement pair is desirable, the complement/anti-complement pair preferably has a binding affinity less than $10^9 \,\mathrm{M}^{-1}$.

An "antibody fragment" is a portion of an antibody such as F(ab')₂, F(ab)₂, Fab', Fab, and the like. Regardless of structure, an antibody fragment binds with the same antigen that is recognized by the intact antibody.

The term "antibody fragment" also includes a synthetic or 55 a genetically engineered polypeptide that binds to a specific antigen, such as polypeptides consisting of the light chain variable region, "Fv" fragments consisting of the variable regions of the heavy and light chains, recombinant single chain polypeptide molecules in which light and heavy vari- 60 able regions are connected by a peptide linker ("scFv proteins"), and minimal recognition units consisting of the amino acid residues that mimic the hypervariable region.

A "chimeric antibody" is a recombinant protein that contains the variable domains and complementary determining 65 regions derived from a rodent antibody, while the remainder of the antibody molecule is derived from a human antibody.

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"Humanized antibodies" are recombinant proteins in which murine complementarity determining regions of a monoclonal antibody have been transferred from heavy and light variable chains of the murine immunoglobulin into a human variable domain.

As used herein, a "therapeutic agent" is a molecule or atom, which is conjugated to an antibody moiety to produce a conjugate, which is useful for therapy. Examples of therapeutic agents include drugs, toxins, immunomodulators, chelators, boron compounds, photoactive agents or dyes, and radioisotopes.

A "detectable label" is a molecule or atom, which can be conjugated to an antibody moiety to produce a molecule useful for diagnosis. Examples of detectable labels include chelators, photoactive agents, radioisotopes, fluorescent agents, paramagnetic ions, or other marker moieties.

The term "affinity tag" is used herein to denote a polypeptide segment that can be attached to a second polypeptide to provide for purification or detection of the second polypeptide or provide sites for attachment of the second polypeptide to a substrate. In principal, any peptide or protein for which an antibody or other specific binding agent is available can be used as an affinity tag. Affinity tags include a poly-histidine tract, protein A (Nilsson et al., EMBO J. 4:1075 (1985); Nilsson et al., Methods Enzymol. 198:3 (1991)), glutathione S transferase (Smith and Johnson, Gene 67:31 (1988)), Glu-Glu affinity tag (Grussenmeyer et al., Proc. Natl. Acad. Sci. USA 82:7952 (1985)), substance P, FLAG peptide (Hopp et al., Biotechnology 6:1204 (1988)), streptavidin binding peptide, or other antigenic epitope or binding domain. See, in general, Ford et al., Protein Expression and Purification 2:95 (1991). DNA molecules encoding affinity tags are available from commercial suppliers (e.g., Pharmacia Biotech, Piscataway, N.J.).

A "naked antibody" is an entire antibody, as opposed to an antibody fragment, which is not conjugated with a therapeutic agent. Naked antibodies include both polyclonal and monoclonal antibodies, as well as certain recombinant antibodies, such as chimeric and humanized antibodies.

As used herein, the term "antibody component" includes both an entire antibody and an antibody fragment.

An "immunoconjugate" is a conjugate of an antibody component with a therapeutic agent or a detectable label.

A "target polypeptide" or a "target peptide" is an amino complement/anti-complement pairs include receptor/ligand 45 acid sequence that comprises at least one epitope, and that is expressed on a target cell, such as a tumor cell, or a cell that carries an infectious agent antigen. T cells recognize peptide epitopes presented by a major histocompatibility complex molecule to a target polypeptide or target peptide and typically lyse the target cell or recruit other immune cells to the site of the target cell, thereby killing the target cell.

An "antigenic peptide" is a peptide, which will bind a major histocompatibility complex molecule to form an MHC-peptide complex, which is recognized by a T cell, thereby inducing a cytotoxic lymphocyte response upon presentation to the T cell. Thus, antigenic peptides are capable of binding to an appropriate major histocompatibility complex molecule and inducing a cytotoxic T cells response, such as cell lysis or specific cytokine release against the target cell, which binds or expresses the antigen. The antigenic peptide can be bound in the context of a class I or class II major histocompatibility complex molecule, on an antigen presenting cell or on a target cell.

In eukaryotes, RNA polymerase II catalyzes the transcription of a structural gene to produce mRNA. A nucleic acid molecule can be designed to contain an RNA polymerase II template in which the RNA transcript has a sequence that is

complementary to that of a specific mRNA. The RNA transcript is termed an "anti-sense RNA" and a nucleic acid molecule that encodes the anti-sense RNA is termed an "antisense gene." Anti-sense RNA molecules are capable of binding to mRNA molecules, resulting in an inhibition of mRNA 5 translation.

Due to the imprecision of standard analytical methods, molecular weights and lengths of polymers are understood to be approximate values. When such a value is expressed as "about" X or "approximately" X, the stated value of X will be 10 understood to be accurate to ±10%.

3. Production of Nucleic Acid Molecules Encoding TACI-Immunoglobulin Proteins

FIG. 1 provides the predicted amino acid sequence of human TACI (von Bülow and Bram, Science 278:138 (1997)). The TACI polypeptide contains the following predicted elements: (a) two cysteine-rich pseudo-repeat structures characteristic of tumor necrosis factor ligand binding 20 domains, (b) a 62 amino acid "stalk region," which resides between the ligand binding domains and the transmembrane domain, (c) a 20 amino acid transmembrane domain, and (d) a 127 amino acid intracellular domain. The amino acid sequence does not contain a predicted hydrophobic amino 25 terminal signal sequence.

In order to create a soluble form of human TACI for use as an inhibitor of the native ligand:native receptor interaction, a TACI extracellular domain—human immunoglobulin Fc fusion protein was generated. The available human TACI 30 sequence was used as the starting point for designing the fusion protein molecule (von Bülow and Bram, Science 278: 138 (1997)). This initial construct, designated as "TACI-Fc4," included amino acid residues 1 through 154 of the TACI polypeptide, and a modified human Fe region, described 35 below. The fusion point of residue 154 was chosen in order to include as much of the stalk region of TACI as possible while not including any potential portion of the predicted transmembrane domain.

Since native TACI polypeptide does not contain an amino 40 terminal signal sequence, an amino terminal signal sequence was added to TACI in order to generate a secreted form of the TACI-Fc fusion protein. The signal sequence was a modified pre-pro sequence from human tissue plasminogen activator. The modifications were included to enhance signal peptidase 45 cleavage and furin protease-specific processing and for that reason this sequence has been referred to as the "optimized tPA (otPA) leader." The otPA sequence (SEQ ID NO:25) is illustrated below; modified amino acid residues are shaded. The recombinant TACI-Fc fusion protein coding sequence 50 was inserted into an expression vector, which was transfected into Chinese hamster ovary cells.

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human IgG Fc antisera revealed two bands, one band was smaller than the expected size of approximately 48 kDa. Amino acid sequence analysis of purified proteins revealed that the smaller band reflected cleavage of TACI fusion proteins at various sites within the TACI stalk region. With reference to SEO ID NO:2, the major termini were found at amino acid residues 118 and 123, although, proteins were also cleaved at amino acid positions 110, 139, and 141.

In addition to heterogeneity caused by cleavage in the stalk region, heterogeneity was also observed at the amino and carboxyl termini. With reference to SEQ ID NO:2, the major amino termini were found at amino acid residues 1, 10, and 13. Differences in the carboxyl terminus reflect the natural heterogeneity of recombinant immunoglobulins and immunoglobulin fusion proteins, which includes the incomplete removal of the carboxyl-terminally-encoded lysine residue. Another source of heterogeneity was found in the variable nature of the carbohydrate structure attached to the Fc encoded immunoglobulin C_{H2} domain.

New versions of TACI-Fc were generated to address the observed heterogeneity. Constructs were designed that included at least one of the following variations in the TACI moiety: (1) portions of the TACI stalk region were deleted, (2) a portion of the TACI stalk region was replaced with a portion of the BCMA stalk region, (3) the arginine residue at position 119 was mutated to eliminate a potential furin cleavage site, (4) the glutamine residue at position 121 was mutated to eliminate a potential furin cleavage site, (5) the arginine residue at position 122 was mutated to eliminate a potential furin cleavage site, (6) amino acid residue at positions 123 and 142 were mutated to amino acid residues found in corresponding positions of murine TACI, (7) the human otPA signal sequence was replaced with a human heavy chain variable region signal sequence, (8) the valine residue at position 29 was mutated to methionine, and the otPA signal sequence was joined in an amino terminal position to this residue, and (9) the otPA signal sequence was joined in an amino terminal location to the alanine residue at position 30.

Modifications were also introduced in the immunoglobulin moiety. Five classes of immunoglobulin, IgG, IgA, IgM, IgD, and IgE, have been identified in higher vertebrates. IgG, IgD, and IgE proteins are characteristically disulfide linked heterotetramers consisting of two identical heavy chains and two identical light chains. Typically, IgM is found as a pentamer of a tetramer, whereas IgA occurs as a dimer of a tetramer.

IgG comprises the major class as it normally exists as the second most abundant protein found in plasma. In humans, IgG consists of four subclasses, designated IgG1, IgG2, IgG3, and IgG4. As shown in FIG. 2, each immunoglobulin heavy chain possesses a constant region that consists of con-

TACI-Fc4 protein at a low level of about 0.3 pg/cell/day. Western blot analysis of TACI-Fc protein with goat anti-

Transfected Chinese hamster ovary cells produced the 65 stant region protein domains (C_{H1} , hinge, C_{H2} , and C_{H3}) that are invariant for a given subclass. The heavy chain constant regions of the IgG class are identified with the Greek symbol

 γ . For example, immunoglobulins of the IgG1 subclass contain a $\gamma 1$ heavy chain constant region.

The Fc fragment, or Fc domain, consists of the disulfide linked heavy chain hinge regions, C_{H2} , and C_{13} domains. In immunoglobulin fusion proteins, Fc domains of the IgG1 5 subclass are often used as the immunoglobulin moiety, because IgG1 has the longest serum half-life of any of the serum proteins. Lengthy serum half-life can be a desirable protein characteristic for animal studies and potential human therapeutic use. In addition, the IgG1 subclass possesses the strongest ability to carry out antibody mediated effector functions. The primary effector function that may be most useful in an immunoglobulin fusion protein is the ability for an IgG1 antibody to mediate antibody dependent cellular cytotoxicity. On the other hand, this could be an undesirable function for a 15 fusion protein that functions primarily as an antagonist. Several of the specific amino acid residues that are important for antibody constant region-mediated activity in the IgG1 subclass have been identified. Inclusion or exclusion of these specific amino acids therefore allows for inclusion or exclusion of specific immunoglobulin constant region-mediated activity.

Six versions of a modified human IgG1 Fc were generated for creating Fc fusion proteins. Fc-488 was designed for convenient cloning of a fusion protein containing the human 25 γ1 Fc region, and it was constructed using the wild-type human immunoglobulin γ1 constant region as a template. Concern about potential deleterious effects due to an unpaired cysteine residue led to the decision to replace the cysteine (amino acid residue 24 of SEQ ID NO:6) that nor- 30 mally disulfide bonds with the immunoglobulin light chain constant region with a serine residue. An additional change was introduced at the codon encoding EU index position 218 (amino acid residue 22 of SEQ ID NO:6) to introduce a BgIII restriction enzyme recognition site for ease of future DNA 35 manipulations. These changes were introduced into the PCR product encoded on the PCR primers. Due to the location of the BgIII site and in order to complete the Fc hinge region, codons for EU index positions 216 and 217 (amino acid residues 20 and 21 of SEQ ID NO:6) were incorporated in the 40 fusion protein partner sequences.

Fc4, Fc5, and Fc6 contain mutations to reduce effector functions mediated by the Fc by reducing FcγRI binding and complement C1q binding. Fc4 contains the same amino acid substitutions that were introduced into Fc-488. Additional 45 amino acid substitutions were introduced to reduce potential Fc mediated effector functions. Specifically, three amino acid substitutions were introduced to reduce FcγRI binding. These are the substitutions at EU index positions 234, 235, and 237 (amino acid residues 38, 39, and 41 of SEQ ID NO:6). Substitutions at these positions have been shown to reduce binding to FcγRI (Duncan et al., *Nature* 332:563 (1988)). These amino acid substitutions may also reduce FcγRIIa binding, as well as FcγRIII binding (Sondermann et al., *Nature* 406:267 (2000); Wines et al., *J Immunol.* 164:5313 (2000)).

Several groups have described the relevance of EU index positions 330 and 331 (amino acid residues 134 and 135 of SEQ ID NO:6) in complement C1q binding and subsequent complement fixation (Canfield and Morrison, *J. Exp. Med.* 173:1483 (1991); Tao et al., *J. Exp. Med.* 178:661 (1993)). 60 Amino acid substitutions at these positions were introduced in Fc4 to reduce complement fixation. The C_{H3} domain of Fc4 is identical to that found in the corresponding wild-type polypeptide, except for the stop codon, which was changed from TGA to TAA to eliminate a potential dam methylation 65 site when the cloned DNA is grown in dam plus strains of *E. coli*.

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In Fc5, the arginine residue at EU index position 218 was mutated back to a lysine, because the BgIII cloning scheme was not used in fusion proteins containing this particular Fe. The remainder of the Fc5 sequence matches the above description for Fc4.

Fc6 is identical to Fc5 except that the carboxyl terminal lysine codon has been eliminated. The C-terminal lysine of mature immunoglobulins is often removed from mature immunoglobulins post-translationally prior to secretion from B-cells, or removed during serum circulation. Consequently, the C-terminal lysine residue is typically not found on circulating antibodies. As in Fc4 and Fc5 above, the stop codon in the Fc6 sequence was changed to TAA.

Fc7 is identical to the wild-type γ1 Fc except for an amino acid substitution at EU index position 297 located in the C_{H2} domain. EU index position Asn-297 (amino acid residue 101 of SEQ ID NO:6) is a site of N-linked carbohydrate attachment. N-linked carbohydrate introduces a potential source of variability in a recombinantly expressed protein due to potential batch-to-batch variations in the carbohydrate structure. In an attempt to eliminate this potential variability, Asn-297 was mutated to a glutamine residue to prevent the attachment of N-linked carbohydrate at that residue position. The carbohydrate at residue 297 is also involved in Fc binding to the FcyRIII (Sondermann et al., Nature 406:267 (2000)). Therefore, removal of the carbohydrate should decrease binding of recombinant Fc7 containing fusion proteins to the FcyRs in general. As above, the stop codon in the Fc7 sequence was mutated to TAA.

Fc8 is identical to the wild-type immunoglobulin γ1 region shown in SEQ ID NO:6, except that the cysteine residue at EU index position 220 (amino acid residue 24 of SEQ ID NO:6) was replaced with a serine residue. This mutation eliminated the cysteine residue that normally disulfide bonds with the immunoglobulin light chain constant region.

Illustrative TACI-Fc constructs are described in Table 1.

TABLE 1

Illustrative TACI-Fc Fusion Protein Constructs						
TACI Sequence ^a	Fc Version					
TACI ^b	Fc4					
$TACI^b$	Fc5					
$TACI^b$	Fcy1					
TACI (d107-154)	Fe5					
TACI (R119Q)	Fc4					
TACI (1-104)-BCMA (42-54) ^c	Fc5					
TACI (d143-150)	Fc5					
TACI (R142G, d143-150)	Fc5					
TACI (R119G, Q121P, R122Q, S123A)	Fc5					
TACI(R119G, R122Q)	Fc5					
TACI (d1-28, V29M)	Fc6					
TACI (d1-29)	Fc6					
TACI (d1-29)	Fc5					
TACI (d1-29, d107-154)	Fc5					
TACI (d1-29, d111-154)	Fc5					
TACI (d1-29, d120-154)	Fc5					

^aInformation about locations, mutations, and deletions of amino acid sequences is provided within parentheses in reference to the amino acid sequence of SEQ ID NO: 2. ^bIncludes amino acid residues 1 to 154 of SEQ ID NO: 2.

The TACI-Fc proteins were produced by recombinant Chinese hamster ovary cells, isolated, and analyzed using Western blot analysis and amino acid sequence analysis. Surprisingly, deletion of the first 29 amino acids from the N-terminus of the TACI polypeptide resulted in a ten-fold increase in the production of TACI-Fc fusion proteins by Chinese hamster ovary cells. This deletion also reduced the cleavage of the

[°]This construct includes amino acid residues 1 to 104 of SEQ ID NO: 2 (TACI) and amino acids 42 to 54 of SEQ ID NO: 27 (BCMA).

full-length stalk region. In addition, cleavage within the TACI stalk region was suppressed either by truncating the TACI stalk region, or by replacing the TACI stalk region within another amino acid sequence (e.g., the amino acid sequence of the BCMA stalk region).

As described in Example 4, functional analyses of TACI-Fc constructs indicate that fusion proteins TACI (d1-29)-Fc5, TACI (d1-29, d107-154)-Fc5, TACI (d1-29, d111-154)-Fc5, and TACI (d1-29, d120-154)-Fc5 have similar binding affinities for ZTNF4. However, constructs, TACI (d1-29)-Fc5, 10 TACI (d1-29, d111-154)-Fc5, and TACI (d1-29, d120-154)-Fc5 appear to bind more ZTNF4 per mole of TACI-Fc than construct, TACI (d1-29, d107-154)-Fc5. Depending upon the intended use (i.e., therapeutic, diagnostic, or research), either high capacity or low capacity TACI-Fc fusion proteins can be 15 employed. In addition, a combination of high capacity and low capacity TACI-Fc fusion proteins enables the titration of ZTNF2 or ZTNF4.

The present invention contemplates TACI-immunoglobulin fusion proteins that comprise a TACI receptor moiety 20 consisting of amino acid residues 30 to 106 of SEQ ID NO:2, 30 to 110 of SEQ ID NO:2, or 30 to 154 of SEQ ID NO:2. The present invention also includes TACI-immunoglobulin fusion proteins that comprise a TACT receptor moiety consisting of amino acid residues 31 to 106 of 25 SEQ ID NO:2, 31 to 110 of SEQ ID NO:2, 31 to 119 of SEQ ID NO:2, or 31 to 154 of SEQ ID NO:2.

More generally, the present invention includes TACI-immunoglobulin fusion proteins, wherein the TACI receptor moiety consists of a fragment of amino acid residues 30 to 30 154 of SEQ ID NO:2, and wherein the TACI receptor moiety binds at least one of ZTNF2 or ZTNF4. Such fragments comprise a cysteine-rich pseudo-repeat region, and optionally, can include at least one of an N-terminal segment, which resides in an amino-terminal position to the cysteine-rich 35 pseudo-repeat region, and a stalk segment, which resides in a carboxyl-terminal position to the cysteine-rich pseudo-repeat region. Suitable cysteine-rich pseudo-repeat regions include polypeptides that: (a) comprise at least one of amino acid residues 34 to 66 of SEQ ID NO:2, and amino acid residues 71 40 to 104 of SEQ ID NO:2, (b) comprise both amino acid residues 34 to 66 of SEQ ID NO:2, and amino acid residues 71 to 104 of SEQ ID NO:2, or (c) comprise amino acid residues 34 to 104 of SEQ ID NO:2.

Suitable N-terminal segments include the following with 45 reference to SEQ ID NO:2: amino acid residue 33, amino acid residues 32 to 33, amino acid residues 31 to 33, and amino acid residues 30 to 33. Suitable stalk segments include one or more amino acids of amino acid residues 105 to 154 of SEQ ID NO:2. For example, the stalk segment can consist of the 50 following with reference to SEQ ID NO:2: amino acid residue 105, amino acid residues 105 to 106, amino acid residues 105 to 107, amino acid residues 105 to 108, amino acid residues 105 to 109, amino acid residues 105 to 110, amino acid residues 105 to 111, amino acid residues 105 to 112, amino 55 acid residues 105 to 113, amino acid residues 105 to 114, amino acid residues 105 to 115, amino acid residues 105 to 116, amino acid residues 105 to 117, amino acid residues 105 to 118, amino acid residues 105 to 119, amino acid residues 105 to 120, amino acid residues 105 to 121, amino acid 60 residues 105 to 122, amino acid residues 105 to 123, amino acid residues 105 to 124, amino acid residues 105 to 125, amino acid residues 105 to 126, amino acid residues 105 to 127, amino acid residues 105 to 128, amino acid residues 105 to 129, amino acid residues 105 to 130, amino acid residues 105 to 131, amino acid residues 105 to 132, amino acid residues 105 to 133, amino acid residues 105 to 134, amino

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acid residues 105 to 135, amino acid residues 105 to 136, amino acid residues 105 to 137, amino acid residues 105 to 138, amino acid residues 105 to 139, amino acid residues 105 to 140, amino acid residues 105 to 141, amino acid residues 105 to 142, amino acid residues 105 to 143, amino acid residues 105 to 144, amino acid residues 105 to 145, amino acid residues 105 to 146, amino acid residues 105 to 147, amino acid residues 105 to 148, amino acid residues 105 to 149, amino acid residues 105 to 150, amino acid residues 105 to 151, amino acid residues 105 to 152, amino acid residues 105 to 153, and amino acid residues 105 to 154.

Additional suitable stalk segments include one or more amino acids of the BCMA stalk region (i.e., amino acid residues 42 to 54 of SEQ ID NO:27. For example, a stalk segment can consist of the following with reference to SEQ ID NO:27: amino acid residue 42, amino acid residues 42 to 43, amino acid residues 42 to 44, amino acid residues 42 to 45, amino acid residues 42 to 46, amino acid residues 42 to 47, amino acid residues 42 to 50, amino acid residues 42 to 51, amino acid residues 42 to 52, amino acid residues 42 to 53, and amino acid residues 42 to 54.

More generally, a stalk segment can consist of two to 50 amino acid residues.

The immunoglobulin moiety of a fusion protein described herein comprises at least one constant region of an immunoglobulin. Preferably, the immunoglobulin moiety represents a segment of a human immunoglobulin. The human immunoglobulin sequence can be a wild-type amino acid sequence, or a modified wild-type amino acid sequence, which has at least one of the amino acid mutations discussed above.

The human immunoglobulin amino acid sequence can also vary from wild-type by having one or more mutations characteristic of a known allotypic determinant. Table 2 shows the allotypic determinants of the human IgG γ 1 constant region (Putman, *The Plasma Proteins, Vol. V*, pages 49 to 140 (Academic Press, Inc. 1987)). EU index positions 214, 356, 358, and 431 define the known IgG γ 1 allotypes. Position 214 is in the C $_{H1}$ domain of the IgG γ 1 constant region, and, therefore, does not reside within the Fc sequence. The wild-type Fc sequence of SEQ ID NO:6 includes the G1m(1) and G1m(2) allotypes. However, the Fc moiety of a TACI-Fc protein can be modified to reflect any combination of these allotypes.

TABLE 2

	Allotypic Determir Immunoglobulin γ		
	Amino Acid	Amino A	Acid Position
Allotype	Residue	EU Index	SEQ ID NO: 6
G1m(1)	Asp, Leu	356, 358	160, 162
G1m(1-)	Glu, Met	356, 358	160, 162
G1m(2)	Gly	431	235
G1m(2-)	Ala	431	235
G1m(3)	Arg	214	_
G1m(3-)	Lys	214	_

The examples of TACI-Fc proteins disclosed herein comprise human IgG1 constant regions. However, suitable immunoglobulin moieties also include polypeptides comprising at least one constant region, such as a heavy chain constant region from any of the following immunoglobulins: IgG2, IgG3, IgG4, IgA1, IgA2, IgD, IgE, and IgM. Advantageously, immunoglobulin moieties derived from wild-type IgG2 or wild-type IgG4 offer reduced effector function, compared with wild-type IgG1 or wild-type IgG3. The present invention

also contemplates fusion proteins that comprise a TACI receptor moiety, as described above, and either albumin or β2-macroglobulin.

Another type of receptor fusion protein that binds ZTNF2 or ZTNF4 is a BCMA-immunoglobulin fusion protein. Studies have been performed with a BCMA-Fc4 fusion protein in which the BCMA moiety consists of amino acid residues 1 to 48 of SEQ ID NO:27. Surprisingly, pharmacokinetic studies in mice revealed that BCMA-Fc4 fusion protein had a halflife of about 101 hours, whereas a TACI-Fc protein had a 10 half-life of 25 hours. Thus, administration of a BCMA-immunoglobulin fusion protein may be preferred in certain clinical settings. Moreover, a combination of TACI-immunoglobulin and BCMA-immunoglobulin fusion proteins may be advantageous to treat certain conditions. This combination therapy can be achieved by administering TACI-immunoglobulin and BCMA-immunoglobulin fusion proteins, or by administering heterodimers of TACI-immunoglobulin and BCMA-immunoglobulin fusion proteins.

Another type of receptor fusion protein that binds ZTNF4 20 is an immunoglobulin fusion protein comprising an extracellular domain of a receptor designated as "Ztnfr12." Ztnfr12 amino acid and nucleotide sequences are provided as SEQ ID NO:59 and SEQ ID NO:60, respectively. Suitable Ztnfr12 acid residues 1 to 69 of SEQ ID NO:60, or amino acid residues 19 to 35 of SEQ ID NO:60.

The fusion proteins of the present invention can have the form of single chain polypeptides, dimers, trimers, or multiples of dimers or trimers. Dimers can be homodimers or 30 heterodimers, and trimers can be homotrimers or heterotrimers. Examples of heterodimers include a TACI-immunoglobulin polypeptide with a BCMA-immunoglobulin polypeptide, a TACI-immunoglobulin polypeptide with a Ztnfr12immunoglobulin polypeptide, and BCMA- 35 a immunoglobulin polypeptide with a Ztnfr12immunoglobulin polypeptide. Examples of heterotrimers include a TACI-immunoglobulin polypeptide with two BCMA-immunoglobulin polypeptides, a TACI-immunoglobulin polypeptide with two Ztnfr12-immunoglobulin 40 polypeptides, a BCMA-immunoglobulin polypeptide with two Ztnfr12-immunoglobulin polypeptides, two TACI-immunoglobulin polypeptides with a BCMA-immunoglobulin polypeptide, two TACI-immunoglobulin polypeptides with a Ztnfr12-immunoglobulin polypeptide, two BCMA-immuno- 45 globulin polypeptides with a Ztnfr12-immunoglobulin polypeptide, and a trimer of a TACI-immunoglobulin polypeptide, a BCMA-immunoglobulin polypeptide, and a Ztnfr12-immunoglobulin polypeptide.

In such fusion proteins, the TACI receptor moiety can 50 comprise at least one of the following amino acid sequences of SEQ ID NO:2: amino acid residues 30 to 154, amino acid residues 34 to 66, amino acid residues 71 to 104, amino acid residues 47 to 62, and amino acid residues 86 to 100. The BCMA receptor moiety can comprise at least one of the 55 following amino acid sequences of SEQ ID NO:27: amino acid residues 1 to 48, amino acid residues 8 to 41, and amino acid residues 21 to 37. The Ztnfr12 receptor moiety can comprise at least one of the following amino acid sequences of SEQ ID NO:60: amino acid residues 1 to 69, and amino 60 acid residues 19 to 35.

Fusion proteins can be produced using the PCR methods used to construct the illustrative TACI-Fc molecules, which are described in the Examples. However, those of skill in the art can use other standard approaches. For example, nucleic 65 acid molecules encoding TACI, BCMA, Ztnfr12, or immunoglobulin polypeptides can be obtained by screening human

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cDNA or genomic libraries using polynucleotide probes based upon sequences disclosed herein. These techniques are standard and well-established (see, for example, Ausubel et al. (eds.), Short Protocols in Molecular Biology, 3rd Edition, pages 4-1 to 4-6 (John Wiley & Sons 1995) ("Ausubel (1995)"); Wu et al., Methods in Gene Biotechnology, pages 33-41 (CRC Press, Inc. 1997) ("Wu (1997)"); Ausubel (1995) at pages 5-1 to 5-6; Wu (1997) at pages 307-327)).

Alternatively, molecules for constructing immunoglobulin fusion proteins can be obtained by synthesizing nucleic acid molecules using mutually priming long oligonucleotides and the nucleotide sequences described herein (see, for example, Ausubel (1995) at pages 8-8 to 8-9). Established techniques using the polymerase chain reaction provide the ability to synthesize DNA molecules at least two kilobases in length (Adang et al., Plant Molec. Biol. 21:1131 (1993), Bambot et al., PCR Methods and Applications 2:266 (1993), Dillon et al., "Use of the Polymerase Chain Reaction for the Rapid Construction of Synthetic Genes," in Methods in Molecular Biology, Vol. 15: PCR Protocols: Current Methods and Applications, White (ed.), pages 263-268, (Humana Press, Inc. 1993), and Holowachuk et al., PCR Methods Appl. 4:299

The nucleic acid molecules of the present invention can receptor moieties include polypeptides comprising amino 25 also be synthesized with "gene machines" using protocols such as the phosphoramidite method. If chemically-synthesized double stranded DNA is required for an application such as the synthesis of a gene or a gene fragment, then each complementary strand is made separately. The production of short genes (60 to 80 base pairs) is technically straightforward and can be accomplished by synthesizing the complementary strands and then annealing them. For the production of longer genes (>300 base pairs), however, special strategies may be required, because the coupling efficiency of each cycle during chemical DNA synthesis is seldom 100%. To overcome this problem, synthetic genes (double-stranded) are assembled in modular form from single-stranded fragments that are from 20 to 100 nucleotides in length. For reviews on polynucleotide synthesis, see, for example, Glick and Pasternak, Molecular Biotechnology, Principles and Applications of Recombinant DNA (ASM Press 1994), Itakura et al., Annu. Rev. Biochem. 53:323 (1984), and Climie et al., Proc. Nat'l Acad. Sci. USA 87:633 (1990).

4. Production of TACI-Immunoglobulin Polypeptides

The polypeptides of the present invention can be produced in recombinant host cells following conventional techniques. To express a TACI-immunoglobulin-encoding sequence, a nucleic acid molecule encoding the polypeptide must be operably linked to regulatory sequences that control transcriptional expression in an expression vector and then, introduced into a host cell. In addition to transcriptional regulatory sequences, such as promoters and enhancers, expression vectors can include translational regulatory sequences and a marker gene, which is suitable for selection of cells that carry the expression vector.

Expression vectors that are suitable for production of a foreign protein in eukaryotic cells typically contain (1) prokaryotic DNA elements coding for a bacterial replication origin and an antibiotic resistance marker to provide for the growth and selection of the expression vector in a bacterial host; (2) eukaryotic DNA elements that control initiation of transcription, such as a promoter; and (3) DNA elements that control the processing of transcripts, such as a transcription termination/polyadenylation sequence.

Expression vectors can also include nucleotide sequences encoding a secretory sequence that directs the heterologous polypeptide into the secretory pathway of a host cell. For example, an expression vector may comprise a nucleotide sequence that encodes TACI-immunoglobulin and a secretory sequence derived from any secreted gene. As discussed above, one suitable signal sequence is a tPA signal sequence. An exemplary tPA signal sequence is provided by SEQ ID NO:25. Another suitable signal sequence is a murine 26-10 V_H signal sequence. The murine 26-10 antibody is described, 10 for example, by Near et al., Mol. Immunol. 27:901 (1990). Illustrative amino acid and nucleotide sequences of a murine $26-10V_H$ signal sequence are provided by SEQ ID NO:61 and SEQ ID NO:65, respectively. SEQ ID NO:62 discloses the amino acid sequence of a TACI-Fc5 fusion protein that com- 15 prises a murine $26-10\,\mathrm{V}_H$ signal sequence.

TACI-immunoglobulin proteins of the present invention may be expressed in mammalian cells. Examples of suitable mammalian host cells include African green monkey kidney cells (Vero; ATCC CRL 1587), human embryonic kidney cells (293-HEK; ATCC CRL 1573), baby hamster kidney cells (BHK-21, BHK-570; ATCC CRL 8544, ATCC CRL 10314), canine kidney cells (MDCK; ATCC CCL 34), Chinese hamster ovary cells (CHO-KI; ATCC CCL61; CHO DG44 (Chasin et al., *Som. Cell. Molec. Genet.* 12:555, 25 1986)), rat pituitary cells (GH1; ATCC CCL82), HeLa S3 cells (ATCC CCL2.2), rat hepatoma cells (H-4-II-E; ATCC CRL 1548) SV40-transformed monkey kidney cells (COS-1; ATCC CRL 1650) and murine embryonic cells (NIH-3T3; ATCC CRL 1658).

For a mammalian host, the transcriptional and translational regulatory signals may be derived from viral sources, such as adenovirus, bovine papilloma virus, simian virus, or the like, in which the regulatory signals are associated with a particular gene which has a high level of expression. Suitable transcriptional and translational regulatory sequences also can be obtained from mammalian genes, such as actin, collagen, myosin, and metallothionein genes.

Transcriptional regulatory sequences include a promoter region sufficient to direct the initiation of RNA synthesis. 40 Suitable eukaryotic promoters include the promoter of the mouse metallothionein I gene (Hamer et al., J. Molec. Appl. Genet. 1:273 (1982)), the TK promoter of Herpes virus (McKnight, Cell 31:355 (1982)), the SV40 early promoter (Benoist et al., Nature 290:304 (1981)), the Rous sarcoma 45 virus promoter (Gorman et al., Proc. Nat'l Acad. Sci. USA 79:6777 (1982)), the cytomegalovirus promoter (Foecking et al., Gene 45:101 (1980)), and the mouse mammary tumor virus promoter (see, generally, Etcheverry, "Expression of Engineered Proteins in Mammalian Cell Culture," in Protein 50 Engineering: Principles and Practice, Cleland et al. (eds.), pages 163-181 (John Wiley & Sons, Inc. 1996)). One useful combination of a promoter and enhancer is provided by a myeloproliferative sarcoma virus promoter and a human cytomegalovirus enhancer.

Alternatively, a prokaryotic promoter, such as the bacteriophage T3 RNA polymerase promoter, can be used to control production of TACI-immunoglobulin proteins in mammalian cells if the prokaryotic promoter is regulated by a eukaryotic promoter (Zhou et al., *Mol. Cell. Biol.* 10:4529 60 (1990), and Kaufman et al., *Nucl. Acids Res.* 19:4485 (1991)).

An expression vector can be introduced into host cells using a variety of standard techniques including calcium phosphate transfection, liposome-mediated transfection, microprojectile-mediated delivery, electroporation, and the 65 like. The transfected cells can be selected and propagated to provide recombinant host cells that comprise the expression

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vector stably integrated in the host cell genome. Techniques for introducing vectors into eukaryotic cells and techniques for selecting such stable transformants using a dominant selectable marker are described, for example, by Ausubel (1995) and by Murray (ed.), *Gene Transfer and Expression Protocols* (Humana Press 1991).

For example, one suitable selectable marker is a gene that provides resistance to the antibiotic neomycin. In this case, selection is carried out in the presence of a neomycin-type drug, such as G-418 or the like. Selection systems can also be used to increase the expression level of the gene of interest, a process referred to as "amplification." Amplification is carried out by culturing transfectants in the presence of a low level of the selective agent and then increasing the amount of selective agent to select for cells that produce high levels of the products of the introduced genes. A suitable amplifiable selectable marker is dihydrofolate reductase, which confers resistance to methotrexate. Other drug resistance genes (e.g., hygromycin resistance, multi-drug resistance, puromycin acetyltransferase) can also be used. Alternatively, markers that introduce an altered phenotype, such as green fluorescent protein, or cell surface proteins such as CD4, CD8, Class I MHC, placental alkaline phosphatase may be used to sort transfected cells from untransfected cells by such means as FACS sorting or magnetic bead separation technology.

TACI-immunoglobulin polypeptides can also be produced by cultured mammalian cells using a viral delivery system. Exemplary viruses for this purpose include adenovirus, herpesvirus, vaccinia virus and adeno-associated virus (AAV). Adenovirus, a double-stranded DNA virus, is currently the best studied gene transfer vector for delivery of heterologous nucleic acid (for a review, see Becker et al., *Meth. Cell Biol.* 43:161 (1994), and Douglas and Curiel, *Science & Medicine* 4:44 (1997)). Advantages of the adenovirus system include the accommodation of relatively large DNA inserts, the ability to grow to high-titer, the ability to infect a broad range of mammalian cell types, and flexibility that allows use with a large number of available vectors containing different promoters.

By deleting portions of the adenovirus genome, larger inserts (up to 7 kb) of heterologous DNA can be accommodated. These inserts can be incorporated into the viral DNA by direct ligation or by homologous recombination with a co-transfected plasmid. An option is to delete the essential E1 gene from the viral vector, which results in the inability to replicate unless the E1 gene is provided by the host cell. Adenovirus vector-infected human 293 cells (ATCC Nos. CRL-1573, 45504, 45505), for example, can be grown as adherent cells or in suspension culture at relatively high cell density to produce significant amounts of protein (see Gamier et al., *Cytotechnol.* 15:145 (1994)).

Those of skill in the art can devise suitable expression vectors for producing the fusion proteins described herein with mammalian cells. Example 4 describes features of one 55 expression vector. As another example, an expression vector can comprise a bicistronic expression cassette that includes a portion of the human cytomegalovirus enhancer, the myeloproliferative sarcoma virus promoter, a nucleotide sequence encoding a fusion protein, the poliovirus internal ribosomal entry sites, a nucleotide sequence encoding murine dihydrofolate reductase, followed by the SV40 poly A addition sequence. The nucleotide sequence of SEQ ID NO:69 shows a cytomegalovirus enhancer/myeloproliferative sarcoma virus LTR promoter construct, in which the cytomegalovirus enhancer extends from nucleotide 1 to 407. The myeloproliferative sarcoma virus LTR promoter, absent the negative control region extends from nucleotide 408 to nucleotide 884 of

SEQ ID NO:69. A nucleotide sequence for the myeloproliferative sarcoma virus LTR promoter without the negative control region is provided in SEQ ID NO:70.

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Example 1 describes an expression vector that comprises a cytomegalovirus promoter to direct the expression of the 5 recombinant protein transgene, an immunoglobulin intron, and a tissue plasminogen activator signal sequence. One suitable immunoglobulin intron is a murine 26-10 V_H intron. SEQ ID NO:66 provides an illustrative nucleotide sequence of a murine 26-10 V_H intron. An expression vector may also 10 include a 5' untranslated region (UTR) located upstream of the nucleotide sequence that encodes a TACI-immunoglobulin protein. A suitable 5'-UTR can be derived from the murine 26-10 V_H gene. SEQ ID NO:63 discloses the nucleotide sequence of a useful native murine 26-10 V_H 5'-UTR, while 15 SEQ ID NO:64 shows the nucleotide sequence of a murine 26-10 V_H 5'-UTR, which has been optimized at the 3' end.

As an illustration, SEQ ID NO:67 provides a nucleotide sequence that includes the following elements: a native murine 26-10 $\rm V_H$ 5'-UTR (nucleotides 1 to 51), a murine 20-10 $\rm V_H$ signal sequence (nucleotides 52 to 97, and 182 to 192), a murine 26-10 $\rm V_H$ intron (nucleotides 98 to 181), a nucleotide sequence that encodes a TACI moiety (nucleotides 193 to 435), and a nucleotide sequence that encodes an Fc5 moiety (nucleotides 436 to 1131). The nucleotide sequence of 25 SEQ ID NO:68 differs from SEQ ID NO:67 due to the replacement of an optimized murine 26-10 $\rm V_H$ 5'-UTR (nucleotides 1 to 51) for the native sequence.

TACI-immunoglobulin proteins can also be expressed in other higher eukaryotic cells, such as avian, fungal, insect, 30 yeast, or plant cells. The baculovirus system provides an efficient means to introduce cloned genes into insect cells. Suitable expression vectors are based upon the Autographa californica multiple nuclear polyhedrosis virus (AcMNPV), and contain well-known promoters such as Drosophila heat 35 shock protein (hsp) 70 promoter, Autographa californica nuclear polyhedrosis virus immediate-early gene promoter (ie-1) and the delayed early 39K promoter, baculovirus p10 promoter, and the Drosophila metallothionein promoter. A second method of making recombinant baculovirus utilizes a 40 transposon-based system described by Luckow (Luckow, et al., J. Virol. 67:4566 (1993)). This system, which utilizes transfer vectors, is sold in the BAC-to-BAC kit (Life Technologies, Rockville, Md.). This system utilizes a transfer vector, PFASTBAC (Life Technologies) containing a Tn7 45 transposon to move the DNA encoding the TACI-immunoglobulin polypeptide into a baculovirus genome maintained in E. coli as a large plasmid called a "bacmid." See, Hill-Perkins and Possee, J. Gen. Virol. 71:971 (1990), Bonning, et al., J. Gen. Virol. 75:1551 (1994), and Chazenbalk, and Rapo- 50 port, J. Biol. Chem. 270:1543 (1995). In addition, transfer vectors can include an in-frame fusion with DNA encoding an epitope tag at the C- or N-terminus of the expressed TACIimmunoglobulin polypeptide, for example, a Glu-Glu epitope tag (Grussenmeyer et al., Proc. Nat'l Acad. Sci. 55 82:7952 (1985)). Using a technique known in the art, a transfer vector containing a nucleotide sequence that encodes a TACI-immunoglobulin protein is transformed into E. coli, and screened for bacmids, which contain an interrupted lacZ gene indicative of recombinant baculovirus. The bacmid 60 DNA containing the recombinant baculovirus genome is then isolated using common techniques.

The illustrative PFASTBAC vector can be modified to a considerable degree. For example, the polyhedrin promoter can be removed and substituted with the baculovirus basic 65 protein promoter (also known as Pcor, p6.9 or MP promoter) which is expressed earlier in the baculovirus infection, and

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has been shown to be advantageous for expressing secreted proteins (see, for example, Hill-Perkins and Possee, *J. Gen. Virol.* 71:971 (1990), Bonning, et al., *J. Gen. Virol.* 75:1551 (1994), and Chazenbalk and Rapoport, *J. Biol. Chem.* 270: 1543 (1995). In such transfer vector constructs, a short or long version of the basic protein promoter can be used. Moreover, transfer vectors can be constructed, with secretory signal sequences derived from insect proteins. For example, a secretory signal sequence from Ecdysteroid Glucosyltransferase (EGT), honey bee Melittin (Invitrogen Corporation; Carlsbad, Calif.), or baculovirus gp67 (PharMingen: San Diego, Calif.) can be used in such constructs.

The recombinant virus or bacmid is used to transfect host cells. Suitable insect host cells include cell lines derived from IPLB-Sf-21, a Spodoptera frugiperda pupal ovarian cell line, such as Sf9 (ATCC CRL 1711), Sf21 AE, and Sf21 (Invitrogen Corporation; San Diego, Calif.), as well as Drosophila Schneider-2 cells, and the HIGH FIVEO cell line (Invitrogen) derived from Trichoplusia ni (U.S. Pat. No. 5,300,435). Commercially available serum-free media can be used to grow and to maintain the cells. Suitable media are Sf900 IITM (Life Technologies) or ESF 921TM (Expression Systems) for the Sf9 cells; and Ex-cellO405TM (JRH Biosciences, Lenexa, Kans.) or Express FiveOTM (Life Technologies) for the *T. ni* cells. When recombinant virus is used, the cells are typically grown up from an inoculation density of approximately 2-5× 10^5 cells to a density of $1-2\times10^6$ cells at which time a recombinant viral stock is added at a multiplicity of infection (MOI) of 0.1 to 10, more typically near 3.

Established techniques for producing recombinant proteins in baculovirus systems are provided by Bailey et al., "Manipulation of Baculovirus Vectors," in *Methods in Molecular Biology, Volume 7: Gene Transfer and Expression Protocols*, Murray (ed.), pages 147-168 (The Humana Press, Inc. 1991), by Patel et al., "The baculovirus expression system," in *DNA Cloning 2: Expression Systems, 2nd Edition*, Glover et al. (eds.), pages 205-244 (Oxford University Press 1995), by Ausubel (1995) at pages 16-37 to 16-57, by Richardson (ed.), *Baculovirus Expression Protocols* (The Humana Press, Inc. 1995), and by Lucknow, "Insect Cell Expression Technology," in *Protein Engineering: Principles and Practice*, Cleland et al. (eds.), pages 183-218 (John Wiley & Sons, Inc. 1996).

Fungal cells, including yeast cells, can also be used to express the genes described herein. Yeast species of particular interest in this regard include Saccharomyces cerevisiae, *Pichia pastoris*, and *Pichia methanolica*. Suitable promoters for expression in yeast include promoters from GAL1 (galactose), PGK (phosphoglycerate kinase), ADH (alcohol dehydrogenase), AOX1 (alcohol oxidase), HIS4 (histidinol dehydrogenase), and the like. Many yeast cloning vectors have been designed and are readily available. A vector can be designed to generate constructs utilizing the necessary elements to carry out homologous recombination in yeast (see, for example, Raymond et al., Bio Techniques 26:134 (1999)). For example, such an expression vector can include URA3 and CEN-ARS (autonomously replicating sequence) sequences required for selection and replication in S. cerevisiae. Other suitable vectors include YIp-based vectors, such as YIp5, YRp vectors, such as YRp17, YEp vectors such as YEp13 and YCp vectors, such as YCp19. Methods for transforming S. cerevisiae cells with exogenous DNA and producing recombinant polypeptides from these cells are disclosed by, for example, Kawasaki, U.S. Pat. No. 4,599,311, Kawasaki et al., U.S. Pat. No. 4,931,373, Brake, U.S. Pat. No. 4,870,008, Welch et al., U.S. Pat. No. 5,037,743, and Murray et al., U.S. Pat. No. 4,845,075. Transformed cells are selected

by phenotype determined by the selectable marker, commonly drug resistance or the ability to grow in the absence of a particular nutrient (e.g., leucine). A suitable vector system for use in *Saccharomyces cerevisiae* is the POT1 vector system disclosed by Kawasaki et al. (U.S. Pat. No. 4,931,373), 5 which allows transformed cells to be selected by growth in glucose-containing media. Additional suitable promoters and terminators for use in yeast include those from glycolytic enzyme genes (see, e.g., Kawasaki, U.S. Pat. No. 4,599,311, Kingsman et al., U.S. Pat. No. 4,615,974, and Bitter, U.S. Pat. No. 4,977,092) and alcohol dehydrogenase genes. See also U.S. Pat. Nos. 4,990,446, 5,063,154, 5,139,936, and 4,661,

Transformation systems for other yeasts, including Hansenula polymorpha, Schizosaccharomyces pombe, 15 Kluyveromyces lactis, Kluyveromyces fragilis, Ustilago maydis, Pichia pastoris, Pichia methanolica, Pichia guillermondii and Candida maltosa are known in the art. See, for example, Gleeson et al., J. Gen. Microbiol. 132:3459 (1986), and Cregg, U.S. Pat. No. 4,882,279. Aspergillus cells may be 20 utilized according to the methods of McKnight et al., U.S. Pat. No. 4,935,349. Methods for transforming Acremonium chrysogenum are disclosed by Sumino et al., U.S. Pat. No. 5,162,228. Methods for transforming Neurospora are disclosed by Lambowitz, U.S. Pat. No. 4,486,533.

For example, the use of *Pichia methanolica* as host for the production of recombinant proteins is disclosed by Raymond, U.S. Pat. No. 5,716,808, Raymond, U.S. Pat. No. 5,736,383, Raymond et al., Yeast 14:11-23 (1998), and in international publication Nos. WO 97/17450, WO 97/17451, WO 30 98/02536, and WO 98/02565. DNA molecules for use in transforming P. methanolica will commonly be prepared as double-stranded, circular plasmids, which are preferably linearized prior to transformation. For polypeptide production in P. methanolica, the promoter and terminator in the plasmid 35 can be that of a P. methanolica gene, such as a P. methanolica alcohol utilization gene (AUG1 or AUG2). Other useful promoters include those of the dihydroxyacetone synthase (DHAS), formate dehydrogenase (FMD), and catalase (CAT) genes. To facilitate integration of the DNA into the host 40 chromosome, it is preferred to have the entire expression segment of the plasmid flanked at both ends by host DNA sequences. A suitable selectable marker for use in Pichia methanolica is a P. methanolica ADE2 gene, which encodes phosphoribosyl-5-aminoimidazole carboxylase (AIRC; EC 4.1.1.21), and which allows ade2 host cells to grow in the absence of adenine. For large-scale, industrial processes where it is desirable to minimize the use of methanol, host cells can be used in which both methanol utilization genes (AUG1 and AUG2) are deleted. For production of secreted 50 proteins, host cells can be deficient in vacuolar protease genes (PEP4 and PRB1). Electroporation is used to facilitate the introduction of a plasmid containing DNA encoding a polypeptide of interest into P. methanolica cells. P. methanolica cells can be transformed by electroporation using an 55 exponentially decaying, pulsed electric field having a field strength of from 2.5 to 4.5 kV/cm, preferably about 3.75 kV/cm, and a time constant (t) of from 1 to 40 milliseconds, most preferably about 20 milliseconds.

Expression vectors can also be introduced into plant protoplasts, intact plant tissues, or isolated plant cells. Methods for introducing expression vectors into plant tissue include the direct infection or co-cultivation of plant tissue with *Agrobacterium tuniefaciens*, microprojectile-mediated delivery, DNA injection, electroporation, and the like. See, for 65 example, Horsch et al., *Science* 227:1229 (1985), Klein et al., *Biotechnology* 10:268 (1992), and Miki et al., "Procedures

for Introducing Foreign DNA into Plants," in *Methods in Plant Molecular Biology and Biotechnology*, Glick et al. (eds.), pages 67-88 (CRC Press, 1993).

Alternatively, TACI-immunoglobulin proteins can be produced in prokaryotic host cells. Suitable promoters that can be used to produce TACI-immunoglobulin polypeptides in a prokaryotic host are well-known to those of skill in the art and include promoters capable of recognizing the T4, T3, Sp6 and T7 polymerases, the P_R and P_L promoters of bacteriophage lambda, the trp, recA, heat shock, lacUV5, tac, lpp-lacSpr, phoA, and lacZ promoters of *E. coli*, promoters of *B. subtilis*, the promoters of the bacteriophages of *Bacillus*, *Streptomyces* promoters, the int promoter of bacteriophage lambda, the bla promoter of pBR322, and the CAT promoter of the chloramphenicol acetyl transferase gene. Prokaryotic promoters have been reviewed by Glick, *J. Ind. Microbiol.* 1:277 (1987), Watson et al., *Molecular Biology of the Gene*, 4th Ed. (Benjamin Cummins 1987), and by Ausubel et al. (1995).

Suitable prokaryotic hosts include *E. coli* and *Bacillus subtilus*. Suitable strains of *E. coli* include BL21(DE3), BL21 (DE3)pLysS, BL21(DE3)pLysE, DH1, DH4I, DH5, DH5I, DH5IF', DH5IMCR, DH10B, DH10B/p3, DH11S, C600, HB101, JM101, JM105, JM109, JM110, K38, RR1, Y1088, Y1089, CSH18, ER1451, and ER1647 (see, for example, Brown (ed.), *Molecular Biology Labfax* (Academic Press 1991)). Suitable strains of *Bacillus subtilus* include BR151, YB886, MI119, MI120, and B170 (see, for example, Hardy, "*Bacillus* Cloning Methods," in *DNA Cloning: A Practical Approach*, Glover (ed.) (IRL Press 1985)).

When expressing a TACI-immunoglobulin protein in bacteria such as E. coli, the polypeptide may be retained in the cytoplasm, typically as insoluble granules, or may be directed to the periplasmic space by a bacterial secretion sequence. In the former case, the cells are lysed, and the granules are recovered and denatured using, for example, guanidine isothiocyanate or urea. The denatured polypeptide can then be refolded and dimerized by diluting the denaturant, such as by dialysis against a solution of urea and a combination of reduced and oxidized glutathione, followed by dialysis against a buffered saline solution. In the latter case, the polypeptide can be recovered from the periplasmic space in a soluble and functional form by disrupting the cells (by, for example, sonication or osmotic shock) to release the contents of the periplasmic space and recovering the protein, thereby obviating the need for denaturation and refolding.

Methods for expressing proteins in prokaryotic hosts are well-known to those of skill in the art (see, for example, Williams et al., "Expression of foreign proteins in *E. coli* using plasmid vectors and purification of specific polyclonal antibodies," in *DNA Cloning 2: Expression Systems, 2nd Edition*, Glover et al. (eds.), page 15 (Oxford University Press 1995), Ward et al., "Genetic Manipulation and Expression of Antibodies," in *Monoclonal Antibodies: Principles and Applications*, page 137 (Wiley-Liss, Inc. 1995), and Georgiou, "Expression of Proteins in Bacteria," in *Protein Engineering: Principles and Practice*, Cleland et al. (eds.), page 101 (John Wiley & Sons, Inc. 1996)).

Standard methods for introducing expression vectors into bacterial, yeast, insect, and plant cells are provided, for example, by Ausubel (1995).

General methods for expressing and recovering foreign protein produced by a mammalian cell system are provided by, for example, Etcheverry, "Expression of Engineered Proteins in Mammalian Cell Culture," in *Protein Engineering: Principles and Practice*, Cleland et al. (eds.), pages 163 (Wiley-Liss, Inc. 1996). Standard techniques for recovering protein produced by a bacterial system is provided by, for

example, Grisshammer et al., "Purification of over-produced proteins from *E. coli* cells," in *DNA Cloning 2: Expression Systems*, 2nd Edition, Glover et al. (eds.), pages 59-92 (Oxford University Press 1995). Established methods for isolating recombinant proteins from a baculovirus system are described by Richardson (ed.), *Baculovirus Expression Protocols* (The Humana Press, Inc. 1995).

As an alternative, polypeptides of the present invention can be synthesized by exclusive solid phase synthesis, partial solid phase methods, fragment condensation or classical solution synthesis. These synthesis methods are well-known to those of skill in the art (see, for example, Merrifield, J. Am. Chem. Soc. 85:2149 (1963), Stewart et al., "Solid Phase Peptide Synthesis" (2nd Edition), (Pierce Chemical Co. 1984), Bayer and Rapp, Chem. Pept. Prot. 3:3 (1986), Atherton et al., 15 Solid Phase Peptide Synthesis: A Practical Approach (IRL Press 1989), Fields and Colowick, "Solid-Phase Peptide Synthesis," Methods in Enzymology Volume 289 (Academic Press 1997), and Lloyd-Williams et al., Chemical Approaches to the Synthesis of Peptides and Proteins (CRC Press, Inc. 20 1997)). Variations in total chemical synthesis strategies, such as "native chemical ligation" and "expressed protein ligation" are also standard (see, for example, Dawson et al., Science 266:776 (1994), Hackeng et al., Proc. Nat'l Acad. Sci. USA 94:7845 (1997), Dawson, Methods Enzymol. 287: 34 (1997), 25 Muir et al, Proc. Nat'l Acad. Sci. USA 95:6705 (1998), and Severinov and Muir, J. Biol. Chem. 273:16205 (1998)).

5. Assays for TACI-Immunoglobulin Fusion Proteins

The function of TACI-immunoglobulin fusion proteins can be examined using a variety of approaches to assess the ability of the fusion proteins to bind ZTNF4 or ZTNF2. As an illustration, Example 4 provides methods for measuring ZTNF4 binding affinity and binding capacity.

Alternatively, TACI-immunoglobulin fusion proteins can be characterized by the ability to inhibit the stimulation of human B cells by soluble ZTNF4, as described by Gross et al., international publication No. WO00/40716. Briefly, human B cells are isolated from peripheral blood mononuclear cells 40 using CD19 magnetic beads and the VarioMacs magnetic separation system (Miltenyi Biotec Auburn, Calif.) according to the manufacturer's instructions. Purified B cells are mixed with soluble ZTNF4 (25 ng/ml) and recombinant human IL-4 (10 ng/ml Pharmingen), and the cells are plated onto round 45 bottom 96 well plates at 1×10⁵ cells per well.

Soluble TACI-immunoglobulin proteins can be diluted from about 5 µg/ml to about 6 ng/ml, and incubated with the B cells for five days, pulsing overnight on day four with 1 µCi ³H-thymidine per well. As a control, TACI-immunoglobulin protein can also be incubated with B cells and IL-4 without ZTNF4. Plates are harvested using Packard plate harvester, and counted using the Packard reader.

This general approach was used to examine three TACI-Fc fusion proteins. Although all fusion proteins inhibited B cell 55 proliferation, constructs TACI (d1-29, d111-154)-Fc5 and TACI (d1-29, d120-154)-Fc5 were more potent than TACI (d1-29, d107-154)-Fc5.

Well-established animal models are available to test in vivo efficacy of TACI-immunoglobulin proteins in certain disease 60 states. For example, TACI-immunoglobulin proteins can be tested in a number of animal models of autoimmune disease, such as MRL-lpr/lpr or NZB×NZW F1 congenic mouse strains, which serve as a model of SLE (systemic lupus erythematosus). Such animal models are known in the art 65 (see, for example, Cohen and Miller (Eds.), *Autoimmune Disease Models: A Guidebook* (Academic Press, Inc. 1994).

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Offspring of a cross between New Zealand Black (NZB) and New Zealand White (NZW) mice develop a spontaneous form of SLE that closely resembles SLE in humans. The offspring mice, known as NZBW begin to develop IgM autoantibodies against T-cells at one month of age, and by five to seven months of age, anti-DNA autoantibodies are the dominant immunoglobulin. Polyclonal B-cell hyperactivity leads to overproduction of autoantibodies. The deposition of these autoantibodies, particularly those directed against single stranded DNA, is associated with the development of glomerulonephritis, which manifests clinically as proteinuria, azotemia, and death from renal failure.

Kidney failure is the leading cause of death in mice affected with spontaneous SLE, and in the NZBW strain, this process is chronic and obliterative. The disease is more rapid and severe in females than males, with mean survival of only 245 days as compared to 406 days for the males. While many of the female mice will be symptomatic (proteinuria) by seven to nine months of age, some can be much younger or older when they develop symptoms. The fatal immune nephritis seen in the NZBW mice is very similar to the glomerulonephritis seen in human SLE, making this spontaneous murine model very attractive for testing of potential SLE therapeutics(Putterman and Naparstek, "Murine Models of Spontaneous Systemic Lupus Erythematosus," in Autoimmune Disease Models: A Guidebook, pages 217-234 (Academic Press, Inc., 1994); Mohan et al., J. Immunol. 154:1470 (1995); and Daikh et al., J Immunol. 159:3104 (1997)).

As described by Gross et al., international publication No. WO00/40716, TACI-immunoglobulin proteins can be administered to NZBW mice to monitor its suppressive effect on B cells over the five-week period when, on average, B-cell autoantibody production is believed to be at high levels in NZBW mice. Briefly, 100 8-week old female (NZB×NZW) F_1 mice can be divided into six groups of 15 mice. Prior to treatment, the mice are monitored once a month for urine protein, and blood is drawn for CBC and serum banking. Serum can be screened for the presence of autoantibodies. Because proteinuria is the hallmark sign of glomerulonephritis, urine protein levels are monitored by dipstick at regular intervals over the course of the study. Treatment can begin when mice are approximately five months of age. The mice receive intraperitoneal injections of vehicle only (phosphate buffered saline) or human TACI-immunoglobulin (control protein) or TACI-immunoglobulin protein (e.g., 20 to 100 µg test protein per dose) three times a week for five weeks.

Blood is collected twice during treatment, and will be collected at least twice following treatment. Urine dipstick values for proteinuria and body weights are determined every two weeks after treatment begins. Blood, urine dipstick value and body weight are collected at the time of euthanasia. The spleen and thymus are divided for fluorescent activated cell sorting analysis and histology. Submandibular salivary glands, mesenteric lymph node chain, liver lobe with gall bladder, cecum and large intestine, stomach, small intestine, pancreas, right kidney, adrenal gland, tongue with trachea and esophagus, heart and lungs are also collected for histology.

Murine models for experimental allergic encephalomyelitis have been used as a tool to investigate both the mechanisms of immune-mediated disease, and methods of potential therapeutic intervention. The model resembles human multiple sclerosis, and produces demyelination as a result of T-cell activation to neuroproteins such as myelin basic protein, or proteolipid protein. Inoculation with antigen leads to induction of CD4+, class II MHC-restricted T-cells (Th1). Changes in the protocol for experimental allergic encephalomyelitis can produce acute, chronic-relapsing, or passive-transfer

variants of the model (Weinberg et al., *J. Immunol.* 162:1818 (1999); Mijaba et al., *Cell. Immunol.* 186:94 (1999); and Glabinski, *Meth. Enzym.* 288:182 (1997)).

Gross et al., international publication No. WO00/40716, describe one approach to evaluating the efficacy of TACI- 5 immunoglobulin proteins in the amelioration of symptoms associated with experimental allergic encephalomyelitis. Briefly, 25 female PLxSJL F1 mice (12 weeks old) are given a subcutaneous injection of 125 μg/mouse of antigen (myelin Proteolipid Protein, PLP, residues 139-151), formulated in 10 complete Freund's Adjuvant. The mice are divided into five groups of five mice. Intraperitoneal injections of pertussis toxin (400 ng) are given on Day 0 and 2. The groups are given a 1x, 10x, or 100x dose of TACI-immunoglobulin protein, one group will receive vehicle only, and one group will receive no treatment. Prevention therapy begins on Day 0, intervention therapy begins on day 7, or at onset of clinical signs. Signs of disease, weight loss, and paralysis manifest in approximately 10 to 14 days, and last for about one week. Animals are assessed daily by collecting body weights and 20 assigning a clinical score to correspond to the extent of their symptoms. Clinical signs of experimental allergic encephalomyelitis appear within 10 to 14 days of inoculation and persist for approximately one week. At the end of the study, all animals are euthanized by gas overdose, and necropsied. 25 The brain and spinal column are collected for histology or frozen for mRNA analysis. Body weight and clinical score data are plotted by individual and by group.

In the collagen-induced arthritis model, mice develop chronic inflammatory arthritis, which closely resembles 30 human rheumatoid arthritis. Since collagen-induced arthritis shares similar immunological and pathological features with rheumatoid arthritis, this makes it an ideal model for screening potential human anti-inflammatory compounds. Another advantage in using the collagen-induced arthritis model is 35 that the mechanisms of pathogenesis are known. The T and B cell epitopes on type II collagen have been identified, and various immunological (delayed-type hypersensitivity and anti-collagen antibody) and inflammatory (cytokines, chemokines, and matrix-degrading enzymes) parameters 40 relating to immune-mediating arthritis have been determined, and can be used to assess test compound efficacy in the models (Wooley, Curr. Opin. Rheum. 3:407 (1999); Williams et al., Immunol. 89:9784 (1992); Myers et al., Life Sci. 61:1861 (1997); and Wang et al., Immunol. 92:8955 (1995)). 45

Gross et al., international publication No. WO00/40716, describe a method for evaluating the efficacy of TACI-immunoglobulin proteins in the amelioration of symptoms associated with collagen-induced arthritis. In brief, eight-week old male DBA/1J mice (Jackson Labs) are divided into groups of 50 five mice/group and are given two subcutaneous injections of 50 to 100 μl of 1 mg/ml collagen (chick or bovine origin), at three week intervals. One control does not receive collagen injections. The first injection is formulated in Complete Freund's Adjuvant, and the second injection is formulated in 55 Incomplete Freund's Adjuvant. TACI-immunoglobulin protein is administered prophylactically at or before the second injection, or after the animal develops a clinical score of two or more that persists at least 24 hours. Animals begin to show symptoms of arthritis following the second collagen injec- 60 tion, usually within two to three weeks. For example, TACI-Fc, a control protein, human IgFc, or phosphate-buffered saline (vehicle) can be administered prophylactically beginning seven days before the second injection (day -7). Proteins can be administered at 100 µg, given three times a week as a 65 200 µl intraperitoneal injection, and continued for four weeks.

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In the collagen-induced arthritis model, the extent of disease is evaluated in each paw using a caliper to measure paw thickness and assigning a clinical score to each paw. For example, a clinical score of "0" indicates a normal mouse, a score of "1" indicates that one or more toes are inflamed, a score of "2" indicates mild paw inflammation, a score of "3" indicates moderate paw inflammation, and a score of "4" indicates severe paw inflammation. Animals are euthanized after the disease as been established for a set period of time, usually seven days. Paws are collected for histology or mRNA analysis, and serum is collected for immunoglobulin and cytokine assays.

Myasthenia gravis is another autoimmune disease for which murine models are available. Myasthenia gravis is a disorder of neuromuscular transmission involving the production of autoantibodies directed against the nicotinic acetylcholine receptor. This disease is acquired or inherited with clinical features including abnormal weakness and fatigue on exertion.

A murine model of myasthenia gravis has been established. (Christadoss et al., "Establishment of a Mouse Model of Myasthenia gravis Which Mimics Human Myasthenia gravid Pathogenesis for Immune Intervention," in Immunobiology of Proteins and Peptides VIII, Atassi and Bixler (Eds.), pages 195-199 (1995)). Experimental autoimmune myasthenia gravis is an antibody mediated disease characterized by the presence of antibodies to acetylcholine receptor. These antibodies destroy the receptor leading to defective neuromuscular electrical impulses, resulting in muscle weakness. In the experimental autoimmune myasthenia gravis model, mice are immunized with the nicotinic acetylcholine receptor. Clinical signs of myasthenia gravis become evident weeks after the second immunization. Experimental autoimmune myasthenia gravis is evaluated by several methods including measuring serum levels of acetylcholine receptor antibodies by radioimmunoassay (Christadoss and Dauphinee, J Immunol. 136:2437 (1986); Lindstrom et al., Methods Enzymol. 74:432 (1981)), measuring muscle acetylcholine receptor, or electromyography (Coligan et al. (Eds.), Protocols in Immunology. Vol. 3, page 15.8.1 (John Wiley & Sons, 1997)).

The effect of TACI-immunoglobulin on experimental autoimmune myasthenia gravis can be determined by administering fusion proteins during ongoing clinical myasthenia gravis in B6 mice. For example, 100 B6 mice are immunized with 20 µg acetylcholine receptor in complete Freund's adjuvant on days 0 and 30. Approximately 40 to 60% of mice will develop moderate (grade 2) to severe (grade 3) clinical myasthenia gravis after the boost with acetylcholine receptor. Mice with grade 2 and 3 clinical disease are divided into three groups (with equal grades of weakness) and weighed (mice with weakness also lose weight, since they have difficulty in consuming food and water) and bled for serum (for pretreatment anti-acetylcholine receptor antibody and isotype level). Group A is injected I.P with phosphate buffered saline, group B is injected intraperitoneally with human IgG-Fc as a control protein (100 µg), and group C is injected with 100 µg of TACI-Fc three times a week for four weeks. Mice are screened for clinical muscle weakness twice a week, and weighed and bled for serum 15 and 30 days after the commencement of treatment. Whole blood is collected on day 15 to determine T/B cell ratio by fluorescence activated cell sorter analysis using markers B220 and CD5. Surviving mice are killed 30 to 45 days after the initiation of treatment, and their carcasses are frozen for later extraction of muscle acetylcholine receptor to determine the loss of muscle acetylcholine receptor, the primary pathology in myasthenia gravis

(see, for example, Coligan et al. (Eds.), *Protocols in Immu-nology. Vol.* 3, page 15.8.1 (John Wiley & Sons, 1997)).

Serum antibodies to mouse muscle acetylcholine receptor can be determined by an established radioimmunoassay, and anti-acetylcholine receptor antibody isotypes (IgM, IgG1, 5 IgG2b and IgG2c) is measured by ELISA. Such methods are known. The effects of TACI-immunoglobulin on ongoing clinical myasthenia gravis, anti-acetylcholine receptor antibody and isotype level, and muscle acetylcholine receptor loss are determined.

Approximately 100 mice can be immunized with 20 μg acetylcholine receptor in complete Freund's adjuvant on day 0 and 30. Mice with clinical myasthenia gravis are divided into four groups. Group A is injected intraperitoneally with 100 μg control Fe, group B is injected with 20 μg control Fc, 15 group C is injected with 100 μg TACI-Fc, and group D is injected with 20 μg TACI-Fc three times a week for four weeks. Mice are weighed and bled for serum before, and 15 and 30 days after the start of the treatment. Serum is tested for anti-acetylcholine receptor antibody and isotypes as 20 described above. Muscle acetylcholine receptor loss can also be measured.

Other suitable assays of TACI-immunoglobulin fusion proteins can be determined by those of skill in the art.

6. Production of TACI-Immunoglobulin Conjugates

The present invention includes chemically modified TACI-immunoglobulin compositions, in which a TACI-immunoglobulin polypeptide is linked with a polymer. Typically, the 30 polymer is water-soluble so that the TACI-immunoglobulin conjugate does not precipitate in an aqueous environment, such as a physiological environment. An example of a suitable polymer is one that has been modified to have a single reactive group, such as an active ester for acylation, or an 35 aldehyde for alkylation, In this way, the degree of polymerization can be controlled. An example of a reactive aldehyde is polyethylene glycol propionaldehyde, or mono-(C₁-C₁₀) alkoxy, or aryloxy derivatives thereof (see, for example, Harris, et al., U.S. Pat. No. 5,252,714). The polymer may be 40 branched or unbranched. Moreover, a mixture of polymers can be used to produce TACI-immunoglobulin conjugates.

TACI-immunoglobulin conjugates used for therapy can comprise pharmaceutically acceptable water-soluble polymer moieties. Suitable water-soluble polymers include polyethylene glycol (PEG), monomethoxy-PEG, mono-(C₁-C₁₀) alkoxy-PEG, aryloxy-PEG, poly-(N-vinyl pyrrolidone)PEG, tresyl monomethoxy PEG, PEG propionaldehyde, bis-succinimidyl carbonate PEG, propylene glycol homopolymers, a polypropylene oxide/ethylene oxide co-polymer, polyoxy-ethylated polyols (e.g., glycerol), polyvinyl alcohol, dextran, cellulose, or other carbohydrate-based polymers. Suitable PEG may have a molecular weight from about 600 to about 60,000, including, for example, 5,000, 12,000, 20,000 and 25,000. A TACI-immunoglobulin conjugate can also comprise a mixture of such water-soluble polymers.

One example of a TACI-immunoglobulin conjugate comprises a TACI-immunoglobulin moiety and a polyalkyl oxide moiety attached to the N-terminus of the TACI-immunoglobulin. PEG is one suitable polyalkyl oxide. As an illustration, 60 TACI-immunoglobulin can be modified with PEG, a process known as "PEGylation." PEGylation of TACI-immunoglobulin can be carried out by any of the PEGylation reactions known in the art (see, for example, EP 0 154 316, Delgado et al., *Critical Reviews in Therapeutic Drug Carrier Systems* 65 9:249 (1992), Duncan and Spreafico, *Clin. Pharmacokinet*. 27:290 (1994), and Francis et al., *Int J Hematol* 68:1 (1998)).

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For example, PEGylation can be performed by an acylation reaction or by an alkylation reaction with a reactive polyethylene glycol molecule. In an alternative approach, TACI-immunoglobulin conjugates are formed by condensing activated PEG, in which a terminal hydroxy or amino group of PEG has been replaced by an activated linker (see, for example, Karasiewicz et al., U.S. Pat. No. 5,382,657).

PEGylation by acylation typically requires reacting an active ester derivative of PEG with a TACI-immunoglobulin polypeptide. An example of an activated PEG ester is PEG esterified to N-hydroxysuccinimide. As used herein, the term "acylation" includes the following types of linkages between TACI-immunoglobulin and a water-soluble polymer: amide, carbamate, urethane, and the like. Methods for preparing PEGylated TACI-immunoglobulin by acylation will typically comprise the steps of (a) reacting a TACI-immunoglobulin polypeptide with PEG (such as a reactive ester of an aldehyde derivative of PEG) under conditions whereby one or more PEG groups attach to TACI-immunoglobulin, and (b) obtaining the reaction product(s). Generally, the optimal reaction conditions for acylation reactions will be determined based upon known parameters and desired results. For example, the larger the ratio of PEG:TACI-immunoglobulin, the greater the percentage of polyPEGylated TACI-immunoglobulin 25 product.

The product of PEGylation by acylation is typically a polyPEGylated TACI-immunoglobulin product, wherein the lysine €-amino groups are PEGylated via an acyl linking group. An example of a connecting linkage is an amide. Typically, the resulting TACI-immunoglobulin will be at least 95% mono-, di-, or tri-pegylated, although some species with higher degrees of PEGylation may be formed depending upon the reaction conditions. PEGylated species can be separated from unconjugated TACI-immunoglobulin polypeptides using standard purification methods, such as dialysis, ultrafiltration, ion exchange chromatography, affinity chromatography, and the like.

PEGylation by alkylation generally involves reacting a terminal aldehyde derivative of PEG with TACI-immunoglobulin in the presence of a reducing agent. PEG groups can be attached to the polypeptide via a —CH₂—NH group.

Derivatization via reductive alkylation to produce a monoPEGylated product takes advantage of the differential reactivity of different types of primary amino groups available for derivatization. Typically, the reaction is performed at a pH that allows one to take advantage of the pKa differences between the ε-amino groups of the lysine residues and the α-amino group of the N-terminal residue of the protein. By such selective derivatization, attachment of a water-soluble polymer that contains a reactive group such as an aldehyde, to a protein is controlled. The conjugation with the polymer occurs predominantly at the N-terminus of the protein without significant modification of other reactive groups such as the lysine side chain amino groups. The present invention provides a substantially homogenous preparation of TACI-immunoglobulin monopolymer conjugates.

Reductive alkylation to produce a substantially homogenous population of monopolymer TACI-immunoglobulin conjugate molecule can comprise the steps of: (a) reacting a TACI-immunoglobulin polypeptide with a reactive PEG under reductive alkylation conditions at a pH suitable to permit selective modification of the α -amino group at the amino terminus of the TACI-immunoglobulin, and (b) obtaining the reaction product(s). The reducing agent used for reductive alkylation should be stable in aqueous solution and able to reduce only the Schiff base formed in the initial process of reductive alkylation. Illustrative reducing agents include

sodium borohydride, sodium cyanoborohydride, dimethylamine borane, trimethylamine borane, and pyridine borane.

For a substantially homogenous population of monopolymer TACI-immunoglobulin conjugates, the reductive alkylation reaction conditions are those that permit the selective attachment of the water soluble polymer moiety to the N-terminus of TACI-immunoglobulin. Such reaction conditions generally provide for pKa differences between the lysine amino groups and the α -amino group at the N-terminus. The pH also affects the ratio of polymer to protein to be used. In general, if the pH is lower, a larger excess of polymer to protein will be desired because the less reactive the N-terminal α -group, the more polymer is needed to achieve optimal conditions. If the pH is higher, the polymer:TACI-immunoglobulin need not be as large because more reactive groups are available. Typically, the pH will fall within the range of 3 to 9, or 3 to 6.

Another factor to consider is the molecular weight of the water-soluble polymer. Generally, the higher the molecular weight of the polymer, the fewer number of polymer molecules which may be attached to the protein. For PEGylation reactions, the typical molecular weight is about 2 kDa to about 100 kDa, about 5 kDa to about 50 kDa, or about 12 kDa to about 25 kDa. The molar ratio of water-soluble polymer to TACT-immunoglobulin will generally be in the range of 1:1 25 to 100:1. Typically, the molar ratio of water-soluble polymer to TACI-immunoglobulin will be 1:1 to 20:1 for polyPEGylation, and 1:1 to 5:1 for monoPEGylation.

General methods for producing conjugates comprising a polypeptide and water-soluble polymer moieties are known ³⁰ in the art. See, for example, Karasiewicz et al., U.S. Pat. No. 5,382,657, Greenwald et al., U.S. Pat. No. 5,738,846, Nieforth et al., *Clin. Pharmaeol. Ther.* 59:636 (1996), Monkarsh et al., *Anal. Biochem.* 247:434 (1997)).

The present invention contemplates compositions comprising a peptide or polypeptide described herein. Such compositions can further comprise a carrier. The carrier can be a conventional organic or inorganic carrier. Examples of carriers include water, buffer solution, alcohol, propylene glycol, macrogol, sesame oil, corn oil, and the like.

7. Isolation of TACI-Immunoglobulin Polypeptides

The polypeptides of the present invention can be purified to at least about 80% purity, to at least about 90% purity, to at 45 least about 95% purity, or greater than 95% purity with respect to contaminating macromolecules, particularly other proteins and nucleic acids, and free of infectious and pyrogenic agents. The polypeptides of the present invention may also be purified to a pharmaceutically pure state, which is 50 greater than 99.9% pure. In certain preparations, purified polypeptide is substantially free of other polypeptides, particularly other polypeptides of animal origin.

Fractionation and/or conventional purification methods can be used to obtain preparations of synthetic TACI-immunoglobulin polypeptides, and recombinant TACI-immunoglobulin polypeptides purified from recombinant host cells. In general, ammonium sulfate precipitation and acid or chaotrope extraction may be used for fractionation of samples. Exemplary purification steps may include hydroxyapatite, 60 size exclusion, FPLC and reverse-phase high performance liquid chromatography. Suitable chromatographic media include derivatized dextrans, agarose, cellulose, polyacrylamide, specialty silicas, and the like. PEI, DEAE, QAE and Q derivatives are suitable. Exemplary chromatographic media include those media derivatized with phenyl, butyl, or octyl groups, such as Phenyl-Sepharose FF (Pharmacia), Toyopearl

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butyl 650 (Toso Haas, Montgomeryville, Pa.), Octyl-Sepharose (Pharmacia) and the like; or polyacrylic resins, such as Amberchrom CG 71 (Toso Haas) and the like. Suitable solid supports include glass beads, silica-based resins, cellulosic resins, agarose beads, cross-linked agarose beads, polystyrene beads, cross-linked polyacrylamide resins and the like that are insoluble under the conditions in which they are to be used. These supports may be modified with reactive groups that allow attachment of proteins by amino groups, carboxyl groups, sulfhydryl groups, hydroxyl groups and/or carbohydrate moieties.

Examples of coupling chemistries include cyanogen bromide activation, N-hydroxysuccinimide activation, epoxide activation, sulfhydryl activation, hydrazide activation, and carboxyl and amino derivatives for carbodiimide coupling chemistries. These and other solid media are well known and widely used in the art, and are available from commercial suppliers. Selection of a particular method for polypeptide isolation and purification is a matter of routine design and is determined in part by the properties of the chosen support. See, for example, *Affinity Chromatography: Principles & Methods* (Pharmacia LKB Biotechnology 1988), and Doonan, *Protein Purification Protocols* (The Humana Press 1996).

Additional variations in TACI-immunoglobulin isolation and purification can be devised by those of skill in the art. For example, anti-TACI or anti-Fc antibodies can be used to isolate large quantities of protein by immunoaffinity purification

The polypeptides of the present invention can also be isolated by exploitation of particular properties. For example, immobilized metal ion adsorption (IMAC) chromatography can be used to purify histidine-rich proteins, including those comprising polyhistidine tags. Briefly, a gel is first charged with divalent metal ions to form a chelate (Sulkowski, *Trends in Biochem.* 3:1 (1985)). Histidine-rich proteins will be adsorbed to this matrix with differing affinities, depending upon the metal ion used, and will be eluted by competitive elution, lowering the pH, or use of strong chelating agents. Other methods of purification include purification of glycosylated proteins by lectin affinity chromatography, Protein A chromatography, and ion exchange chromatography (M. Deutscher, (ed.), *Meth. Enzymol.* 182:529 (1990)).

TACI-immunoglobulin polypeptides or fragments thereof may also be prepared through chemical synthesis, as described above. TACI-immunoglobulin polypeptides may be monomers or multimers; glycosylated or non-glycosylated; PEGylated or non-PEGylated; and may or may not include an initial methionine amino acid residue. A TACI-immunoglobulin fusion protein may be non-glycosylated, glycosylated, or glycosylated only in the TACI moiety or in the immunoglobulin moiety. The immunoglobulin moiety can be obtained from a human antibody, a chimeric antibody, or a humanized antibody.

8. Therapeutic Uses of TACI-Immunoglobulin Polypeptides

TACI-immunoglobulin proteins can be used to modulate the immune system by binding ZTNF4 or ZTNF2, and thus, preventing the binding of these ligands with endogenous TACI or BCMA receptors. Accordingly, the present invention includes the use of TACI-immunoglobulin proteins to a subject, which lacks an adequate amount of TACI or BCMA receptors, or which produces an excess of ZTNF4 or ZTNF2. These molecules can be administered to any subject in need of treatment, and the present invention contemplates both vet-

erinary and human therapeutic uses. Illustrative subjects include mammalian subjects, such as farm animals, domestic animals, and human patients.

TACI-immunoglobulin polypeptides can be used for the treatment of autoimmune diseases, B cell cancers, immunomodulation, IBD and any antibody-mediated pathologies (e.g., ITCP, myasthenia gravis and the like), renal diseases, indirect T cell immune response, graft rejection, and graft versus host disease. The polypeptides of the present invention can be targeted to specifically regulate B cell responses during the immune response. Additionally, the polypeptides of the present invention can be used to modulate B cell development, development of other cells, antibody production, and cytokine production. Polypeptides of the present invention can also modulate T and B cell communication by neutralizing the proliferative effects of ZTNF4.

TACI-immunoglobulin polypeptides of the present invention can be useful to neutralize the effects of ZTNF4 for treating pre-B or B-cell leukemias, such as plasma cell leukemia, chronic or acute lymphocytic leukemia, myelomas such as multiple myeloma, plasma cell myeloma, endothelial myeloma and giant cell myeloma, and lymphomas such as non-Hodgkins lymphoma, for which an increase in ZTNF4 polypeptides is associated.

ZTNF4 is expressed in CD8+ cells, monocytes, dendritic cells, activated monocytes, which indicates that, in certain autoimmune disorders, cytotoxic T-cells might stimulate B-cell production through excess production of ZTNF4. Immunosuppressant proteins that selectively block the action 30 of B-Lymphocytes would be of use in treating disease. Autoantibody production is common to several autoimmune diseases and contributes to tissue destruction and exacerbation of disease. Autoantibodies can also lead to the occurrence of immune complex deposition complications and lead to 35 many symptoms of systemic lupus erythematosus, including kidney failure, neuralgic symptoms and death. Modulating antibody production independent of cellular response would also be beneficial in many disease states. B cells have also been shown to play a role in the secretion of arthritogenic 40 immunoglobulins in rheumatoid arthritis. As such, inhibition of ZTNF4 antibody production would be beneficial in treatment of autoimmune diseases such as myasthenia gravis, rheumatoid arthritis, polyarticular-course juvenile rheumatoid arthritis, and psoriatic arthritis. Immunosuppressant 45 therapeutics such as TACI-immunoglobulin proteins that selectively block or neutralize the action of B-lymphocytes would be useful for such purposes.

The invention provides methods employing TACI-immunoglobulin proteins for selectively blocking or neutralizing 50 the actions of B-cells in association with end stage renal diseases, which may or may not be associated with autoimmune diseases. Such methods would also be useful for treating immunologic renal diseases. Such methods would be would be useful for treating glomerulonephritis associated 55 with diseases such as membranous nephropathy, IgA nephropathy or Berger's Disease, IgM nephropathy, Goodpasture's Disease, post-infectious glomerulonephritis, mesangioproliferative disease, chronic lymphoid leukemia, minimal-change nephrotic syndrome. Such methods would also serve as thera- 60 peutic applications for treating secondary glomerulonephritis or vasculitis associated with such diseases as lupus, polyarteritis, Henoch-Schonlein, Scleroderma, HIV-related diseases, amyloidosis or hemolytic uremic syndrome. The methods of the present invention would also be useful as part of a 65 therapeutic application for treating interstitial nephritis or pyelonephritis associated with chronic pyelonephritis, anal34

gesic abuse, nephrocalcinosis, nephropathy caused by other agents, nephrolithiasis, or chronic or acute interstitial nephritis

The methods of the present invention also include use of TACI-immunoglobulin proteins in the treatment of hypertensive or large vessel diseases, including renal artery stenosis or occlusion and cholesterol emboli or renal emboli.

The present invention also provides methods for treatment of renal or urological neoplasms, multiple myelomas, lymphomas, light chain neuropathy or amyloidosis.

The invention also provides methods for blocking or inhibiting activated B cells using TACI-immunoglobulin proteins for the treatment of asthma and other chronic airway diseases such as bronchitis and emphysema. The TACI-immunoglobulin proteins described herein can also be used to treat Sjögren's Syndrome.

Also provided are methods for inhibiting or neutralizing an effector T cell response using TACI-immunoglobulin proteins for use in immunosuppression, in particular for such therapeutic use as for graft-versus-host disease and graft rejection. Moreover, TACI-immunoglobulin proteins would be useful in therapeutic protocols for treatment of such autoimmune diseases as insulin dependent diabetes mellitus (IDDM) and Crohn's Disease. Methods of the present invention would have additional therapeutic value for treating chronic inflammatory diseases, in particular to lessen joint pain, swelling, anemia and other associated symptoms as well as treating septic shock.

Well established animal models are available to test in vivo efficacy of TACI-immunoglobulin proteins of the present invention in certain disease states. In particular, TACI-immunoglobulin proteins can be tested in vivo in a number of animal models of autoimmune disease, such as MRL-lpr/lpr or NZB×NZW F1 congenic mouse strains which serve as a model of SLE (systemic lupus erythematosus). Such animal models are known in the art.

Offspring of a cross between New Zealand Black (NZB) and New Zealand White (NZW) mice develop a spontaneous form of SLE that closely resembles SLE in humans. The offspring mice, known as NZBW begin to develop IgM autoantibodies against T-cells at 1 month of age, and by 5-7 months of age, Ig anti-DNA autoantibodies are the dominant immunoglobulin. Polyclonal B-cell hyperactivity leads to overproduction of autoantibodies. The deposition of these autoantibodies, particularly ones directed against single stranded DNA is associated with the development of glomerulonephritis, which manifests clinically as proteinuria, azotemia, and death from renal failure. Kidney failure is the leading cause of death in mice affected with spontaneous SLE, and in the NZBW strain, this process is chronic and obliterative. The disease is more rapid and severe in females than males, with mean survival of only 245 days as compared to 406 days for the males. While many of the female mice will be symptomatic (proteinuria) by 7-9 months of age, some can be much younger or older when they develop symptoms. The fatal immune nephritis seen in the NZBW mice is very similar to the glomerulonephritis seen in human SLE, making this spontaneous murine model useful for testing of potential SLE therapeutics.

Mouse models for experimental allergic encephalomyelitis (EAE) has been used as a tool to investigate both the mechanisms of immune-mediated disease, and methods of potential therapeutic intervention. The model resembles human multiple sclerosis, and produces demyelination as a result of T-cell activation to neuroproteins such as myelin basic protein (MBP), or proteolipid protein (PLP). Inoculation with antigen leads to induction of CD4+, class II MHC-restricted

T-cells (Th1). Changes in the protocol for EAE can produce acute, chronic-relapsing, or passive-transfer variants of the model.

In the collagen-induced arthritis (CIA) model, mice develop chronic inflammatory arthritis, which closely 5 resembles human rheumatoid arthritis (RA). Since CIA shares similar immunological and pathological features with RA, this makes it an ideal model for screening potential human anti-inflammatory compounds. Another advantage in using the CIA model is that the mechanisms of pathogenesis are known. The T and B cell epitopes on type II collagen have been identified, and various immunological (delayed-type hypersensitivity and anti-collagen antibody) and inflammatory (cytokines, chemokines, and matrix-degrading enzymes) parameters relating to immune-mediating arthritis have been 15 determined, and can be used to assess test compound efficacy in the models.

Myasthenia gravis (MG) is another autoimmune disease for which murine models are available. MG is a disorder of neuromuscular transmission involving the production of 20 autoantibodies directed against the nicotinic acetylcholine receptor (AChR). MG is acquired or inherited with clinical features including abnormal weakness and fatigue on exertion. A mouse model of MG has been established. Experimental autoimmune myasthenia gravis (EAMG) is an anti- 25 body mediated disease characterized by the presence of antibodies to AChR. These antibodies destroy the receptor leading to defective neuromuscular electrical impulses, resulting in muscle weakness. In the EAMG model, mice are immunized with the nicotinic acetylcholine receptor. Clinical 30 signs of MG become evident weeks after the second immunization. EAMG is evaluated by several methods including measuring serum levels of AChR antibodies by radioimmunoassay, measuring muscle AChR, or electromyography.

Generally, the dosage of administered TACI-immunoglobulin protein will vary depending upon such factors as the subject's age, weight, height, sex, general medical condition and previous medical history. Typically, it is desirable to provide the recipient with a dosage of TACI-immunoglobulin protein, which is in the range of from about 1 pg/kg to 10 40 mg/kg (amount of agent/body weight of subject), although a lower or higher dosage also may be administered as circumstances dictate.

Administration of a TACI-immunoglobulin protein to a subject can be intravenous, intraarterial, intraperitoneal, 45 intramuscular, subcutaneous, intrapleural, intrathecal, by perfusion through a regional catheter, or by direct intralesional injection. When administering therapeutic proteins by injection, the administration may be by continuous infusion or by single or multiple boluses.

Additional routes of administration include oral, mucosalmembrane, pulmonary, and transcutaneous. Oral delivery is suitable for polyester microspheres, zein microspheres, proteinoid microspheres, polycyanoacrylate microspheres, and lipid-based systems (see, for example, DiBase and Morrel, 55 "Oral Delivery of Microencapsulated Proteins," in Protein Delivery: Physical Systems, Sanders and Hendren (eds.), pages 255-288 (Plenum Press 1997)). The feasibility of an intranasal delivery is exemplified by such a mode of insulin administration (see, for example, Hinchcliffe and Ilium, Adv. 60 Drug Deliv. Rev. 35:199 (1999)). Dry or liquid particles comprising TACI-immunoglobulin can be prepared and inhaled with the aid of dry-powder dispersers, liquid aerosol generators, or nebulizers (e.g., Pettit and Gombotz, TIBTECH 16:343 (1998); Patton et al., Adv. Drug Deliv. Rev. 35:235 65 (1999)). This approach is illustrated by the AERX diabetes management system, which is a hand-held electronic inhaler

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that delivers aerosolized insulin into the lungs. Studies have shown that proteins as large as 48,000 kDa have been delivered across skin at therapeutic concentrations with the aid of low-frequency ultrasound, which illustrates the feasibility of transcutaneous administration (Mitragotri et al., *Science* 269: 850 (1995)). Transdermal delivery using electroporation provides another means to administer a TACI-immunoglobulin protein (Potts et al., *Pharm. Biotechnol.* 10:213 (1997)).

A pharmaceutical composition comprising a TACI-immunoglobulin protein can be formulated according to known methods to prepare pharmaceutically useful compositions, whereby the therapeutic proteins are combined in a mixture with a pharmaceutically acceptable carrier. A composition is said to be a "pharmaceutically acceptable carrier" if its administration can be tolerated by a recipient patient. Sterile phosphate-buffered saline is one example of a pharmaceutically acceptable carrier. Other suitable carriers are well-known to those in the art. See, for example, Gennaro (ed.), *Remington's Pharmaceutical Sciences*, 19th Edition (Mack Publishing Company 1995).

For purposes of therapy, TACI-immunoglobulin proteins are administered to a patient in a therapeutically effective amount. A TACI-immunoglobulin protein and a pharmaceutically acceptable carrier is said to be administered in a "therapeutically effective amount" if the amount administered is physiologically significant. An agent is physiologically significant if its presence results in a detectable change in the physiology of a recipient patient. For example, an agent used to treat inflammation is physiologically significant if its presence alleviates the inflammatory response. As another example, an agent used to inhibit the growth of tumor cells is physiologically significant if the administration of the agent results in a decrease in the number of tumor cells, decreased metastasis, a decrease in the size of a solid tumor, or increased necrosis of a tumor. Furthermore, an agent used to treat systemic lupus erythematosus is physiologically significant if the administration of the agent results in a decrease of circulating anti-double stranded DNA antibodies, or a decrease in at least one of the following symptoms: fever, joint pain, erythematosus skin lesions, or other features of systemic lupus erythematosus. One example of a general indication that a TACI-immunoglobulin protein is administered in a therapeutically effective amount is that, following administration to a subject, there is a decrease in circulating levels of ZTNF4 (BLyS).

A pharmaceutical composition comprising a TACI-immunoglobulin protein can be furnished in liquid form, in an aerosol, or in solid form. Liquid forms, are illustrated by injectable solutions and oral suspensions. Exemplary solid forms include capsules, tablets, and controlled-release forms. The latter form is illustrated by miniosmotic pumps and implants (Bremer et al., *Pharm. Biotechnol.* 10:239 (1997); Ranade, "Implants in Drug Delivery," in *Drug Delivery Systems*, Ranade and Hollinger (eds.), pages 95-123 (CRC Press 1995); Bremer et al., "Protein Delivery with Infusion Pumps," in *Protein Delivery: Physical Systems*, Sanders and Hendren (eds.), pages 239-254 (Plenum Press 1997); Yewey et al., "Delivery of Proteins from a Controlled Release Injectable Implant," in *Protein Delivery: Physical Systems*, Sanders and Hendren (eds.), pages 93-117 (Plenum Press 1997)).

Liposomes provide one means to deliver therapeutic polypeptides to a subject intravenously, intraperitoneally, intrathecally, intramuscularly, subcutaneously, or via oral administration, inhalation, or intranasal administration. Liposomes are microscopic vesicles that consist of one or more lipid bilayers surrounding aqueous compartments (see, generally, Bakker-Woudenberg et al., Eur. J. Clin. Microbiol.

Infect. Dis. 12 (Suppl. 1):S61 (1993), Kim, Drugs 46:618 (1993), and Ranade, "Site-Specific Drug Delivery Using Liposomes as Carriers," in *Drug Delivery Systems*, Ranade and Hollinger (eds.), pages 3-24 (CRC Press 1995)). Liposomes are similar in composition to cellular membranes and 5 as a result, liposomes can be administered safely and are biodegradable. Depending on the method of preparation, liposomes may be unilamellar or multilamellar, and liposomes can vary in size with diameters ranging from 0.02 μm to greater than 10 µm. A variety of agents can be encapsulated 10 in liposomes: hydrophobic agents partition in the bilayers and hydrophilic agents partition within the inner aqueous space(s) (see, for example, Machy et al., Liposomes In Cell Biology And Pharmacology (John Libbey 1987), and Ostro et al., American J. Hosp. Pharm. 46:1576 (1989)). Moreover, it is 15 possible to control the therapeutic availability of the encapsulated agent by varying liposome size, the number of bilayers, lipid composition, as well as the charge and surface characteristics of the liposomes.

Liposomes can adsorb to virtually any type of cell and then 20 slowly release the encapsulated agent. Alternatively, an absorbed liposome may be endocytosed by cells that are phagocytic. Endocytosis is followed by intralysosomal degradation of liposomal lipids and release of the encapsulated agents (Scherphof et al., *Ann. N.Y. Acad. Sci.* 446:368 25 (1985)). After intravenous administration, small liposomes (0.1 to 1.0 µm) are typically taken up by cells of the reticuloendothelial system, located principally in the liver and spleen, whereas liposomes larger than 3.0 µm are deposited in the lung. This preferential uptake of smaller liposomes by the 30 cells of the reticuloendothelial system has been used to deliver chemotherapeutic agents to macrophages and to tumors of the liver.

The reticuloendothelial system can be circumvented by several methods including saturation with large doses of liposome particles, or selective macrophage inactivation by pharmacological means (Claassen et al., *Biochim. Biophys. Acta* 802:428 (1984)). In addition, incorporation of glycolipid- or polyethelene glycol-derivatized phospholipids into liposome membranes has been shown to result in a significantly 40 reduced uptake by the reticuloendothelial system (Allen et al., *Biochim. Biophys. Acta* 1068:133 (1991); Allen et al., *Biochim. Biophys. Acta* 1150:9 (1993)).

Liposomes can also be prepared to target particular cells or organs by varying phospholipid composition or by inserting 45 receptors or ligands into the liposomes. For example, liposomes, prepared with a high content of a nonionic surfactant, have been used to target the liver (Hayakawa et al., Japanese Patent 04-244,018; Kato et al., *Biol. Pharm. Bull.* 16:960 (1993)). These formulations were prepared by mixing soybean phospatidylcholine, α-tocopherol, and ethoxylated hydrogenated castor oil (HCO-60) in methanol, concentrating the mixture under vacuum, and then reconstituting the mixture with water. A liposomal formulation of dipalmitolylphosphatidylcholine (DPPC) with a soybean-derived sterylglucoside mixture (SG) and cholesterol (Ch) has also been shown to target the liver (Shimizu et al., *Biol. Pharm. Bull.* 20:881 (1997)).

Alternatively, various targeting ligands can be bound to the surface of the liposome, such as antibodies, antibody fragments, carbohydrates, vitamins, and transport proteins. For example, liposomes can be modified with branched type galactosyllipid derivatives to target asialoglycoprotein (galactose) receptors, which are exclusively expressed on the surface of liver cells (Kato and Sugiyama, *Crit. Rev. Ther. 65 Drug Carrier Syst.* 14:287 (1997); Murahashi et al., *Biol. Pharm. Bull.* 20:259 (1997)). Similarly, Wu et al., *Hepatology*

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27:772 (1998), have shown that labeling liposomes with asialofetuin led to a shortened liposome plasma half-life and greatly enhanced uptake of asialofetuin-labeled liposome by hepatocytes. On the other hand, hepatic accumulation of liposomes comprising branched type galactosyllipid derivatives can be inhibited by preinjection of asialofetuin (Murahashi et al., *Biol. Pharm. Bull.* 20:259 (1997)). Polyaconitylated human serum albumin liposomes provide another approach for targeting liposomes to liver cells (Kamps et al., *Proc. Nat'l Acad. Sci. USA* 94:11681 (1997)). Moreover, Geho, et al. U.S. Pat. No. 4,603,044, describe a hepatocyte-directed liposome vesicle delivery system, which has specificity for hepatobiliary receptors associated with the specialized metabolic cells of the liver.

In a more general approach to tissue targeting, target cells are prelabeled with biotinylated antibodies specific for a ligand expressed by the target cell (Harasym et al., *Adv. Drug Deliv. Rev.* 32:99 (1998)). After plasma elimination of free antibody, streptavidin-conjugated liposomes are administered. In another approach, targeting antibodies are directly attached to liposomes (Harasym et al., *Adv. Drug Deliv. Rev.* 32:99 (1998)).

TACI-immunoglobulin proteins can be encapsulated within liposomes using standard techniques of protein microencapsulation (see, for example, Anderson et al., *Infect. Immun.* 31:1099 (1981), Anderson et al., *Cancer Res.* 50:1853 (1990), and Cohen et al., *Biochim. Biophys. Acta* 1063:95 (1991), Alving et al. "Preparation and Use of Liposomes in Immunological Studies," in *Liposome Technology*, 2nd Edition, Vol. III, Gregoriadis (ed.), page 317 (CRC Press 1993), Wassef et al., *Meth. Enzymol.* 149:124 (1987)). As noted above, therapeutically useful liposomes may contain a variety of components. For example, liposomes may comprise lipid derivatives of poly(ethylene glycol) (Allen et al., *Biochim. Biophys. Acta* 1150:9 (1993)).

Degradable polymer microspheres have been designed to maintain high systemic levels of therapeutic proteins. Microspheres are prepared from degradable polymers such as poly (lactide-co-glycolide) (PLG), polyanhydrides, poly (ortho esters), nonbiodegradable ethylvinyl acetate polymers, in which proteins are entrapped in the polymer (Gombotz and Pettit, Bioconjugate Chem. 6:332 (1995); Ranade, "Role of Polymers in Drug Delivery," in Drug Delivery Systems, Ranade and Hollinger (eds.), pages 51-93 (CRC Press 1995); Roskos and Maskiewicz, "Degradable Controlled Release Systems Useful for Protein Delivery," in Protein Delivery: Physical Systems, Sanders and Hendren (eds.), pages 45-92 (Plenum Press 1997); Bartus et al., Science 281:1161 (1998); Putney and Burke, Nature Biotechnology 16:153 (1998); Putney, Curr. Opin. Chem. Biol. 2:548 (1998)). Polyethylene glycol (PEG)-coated nanospheres can also provide carriers for intravenous administration of therapeutic proteins (see, for example, Gref et al., Pharm. Biotechnol. 10:167 (1997)).

The present invention also contemplates chemically modified TACI-immunoglobulin proteins in which the polypeptide is linked with a polymer, as discussed above.

Other dosage forms can be devised by those skilled in the art, as shown, for example, by Ansel and Popovich, *Pharmaceutical Dosage Forms and Drug Delivery Systems*, 5th Edition (Lea & Febiger 1990), Gennaro (ed.), *Remington's Pharmaceutical Sciences*, 19th Edition (Mack Publishing Company 1995), and by Ranade and Hollinger, *Drug Delivery Systems* (CRC Press 1996).

As an illustration, pharmaceutical compositions may be supplied as a kit comprising a container that comprises a TACI-immunoglobulin protein. Therapeutic polypeptides can be provided in the form of an injectable solution for single

or multiple doses, or as a sterile powder that will be reconstituted before injection. Alternatively, such a kit can include a dry-powder disperser, liquid aerosol generator, or nebulizer for administration of a therapeutic polypeptide. Such a kit may further comprise written information on indications and usage of the pharmaceutical composition. Moreover, such information may include a statement that the TACI-immunoglobulin protein composition is contraindicated in patients with known hypersensitivity to either the TACI receptor moiety or the immunoglobulin moiety.

9. Therapeutic Uses of TACI-Immunoglobulin Nucleotide Sequences

The present invention includes the use of nucleic acid 15 molecules that encode TACI-immunoglobulin fusion proteins to provide these fusion proteins to a subject in need of such treatment. For veterinary therapeutic use or human therapeutic use, such nucleic acid molecules can be administered to a subject having a disorder or disease, as discussed 20 above. As one example discussed earlier, nucleic acid molecules encoding a TACI-immunoglobulin fusion protein can be used for long-term treatment of systemic lupus erythematosus.

There are numerous approaches for introducing a TACI- 25 immunoglobulin gene to a subject, including the use of recombinant host cells that express TACI-immunoglobulin, delivery of naked nucleic acid encoding TACI-immunoglobulin, use of a cationic lipid carrier with a nucleic acid molecule that encodes TACI-immunoglobulin, and the use of 30 viruses that express TACI-immunoglobulin, such as recombinant retroviruses, recombinant adeno-associated viruses, recombinant adenoviruses, and recombinant Herpes simplex viruses (see, for example, Mulligan, Science 260:926 (1993), Rosenberg et al., Science 242:1575 (1988), LaSalle et al., 35 Science 259:988 (1993), Wolff et al., Science 247:1465 (1990), Breakfield and Deluca, The New Biologist 3:203 (1991)). In an ex vivo approach, for example, cells are isolated from a subject, transfected with a vector that expresses a TACI-immunoglobulin gene, and then transplanted into the 40 subject.

In order to effect expression of a TACI-immunoglobulin gene, an expression vector is constructed in which a nucleotide sequence encoding a TACI-immunoglobulin gene is operably linked to a core promoter, and optionally a regulatory element, to control gene transcription. The general requirements of an expression vector are described above.

Alternatively, a TACI-immunoglobulin gene can be delivered using recombinant viral vectors, including for example, adenoviral vectors (e.g., Kass-Eisler et al., Proc. Nat'l Acad. 50 Sci. USA 90:11498 (1993), Kolls et al, Proc. Nat'l Acad. Sci. USA 91:215 (1994), Li et al., Hum. Gene Ther. 4:403 (1993), Vincent et al., Nat. Genet. 5:130 (1993), and Zabner et al., Cell 75:207 (1993)), adenovirus-associated viral vectors (Flotte et al., Proc. Nat'l Acad. Sci. USA 90:10613 (1993)), 55 alphaviruses such as Semliki Forest Virus and Sindbis Virus (Hertz and Huang, J. Vir. 66:857 (1992), Raju and Huang, J. Vir. 65:2501 (1991), and Xiong et al., Science 243:1188 (1989)), herpes viral vectors (e.g., U.S. Pat. Nos. 4,769,331, 4,859,587, 5,288,641 and 5,328,688), parvovirus vectors 60 (Koering et al., Hum. Gene Therap. 5:457 (1994)), pox virus vectors (Ozaki et al., Biochem. Biophys. Res. Comm. 193:653 (1993), Panicali and Paoletti, Proc. Nat'l Acad. Sci. USA 79:4927 (1982)), pox viruses, such as canary pox virus or vaccinia virus (Fisher-Hoch et al., Proc. Nat'l Acad. Sci. USA 65 86:317 (1989), and Flexner et al., Ann. N.Y. Acad. Sci. 569:86 (1989)), and retroviruses (e.g., Baba et al., J. Neurosurg

79:729 (1993), Ram et al., *Cancer Res.* 53:83 (1993), Takamiya et al., *J. Neurosci. Res.* 33:493 (1992), Vile and Hart, *Cancer Res.* 53:962 (1993), Vile and Hart, *Cancer Res.* 53:3860 (1993), and Anderson et al., U.S. Pat. No. 5,399, 346). Within various embodiments, either the viral vector itself, or a viral particle, which contains the viral vector may be utilized in the methods and compositions described below.

As an illustration of one system, adenovirus, a double-stranded DNA virus, is a well-characterized gene transfer vector for delivery of a heterologous nucleic acid molecule (for a review, see Becker et al., *Meth. Cell Biol.* 43:161 (1994); Douglas and Curiel, *Science & Medicine* 4:44 (1997)). The adenovirus system offers several advantages including: (i) the ability to accommodate relatively large DNA inserts, (ii) the ability to be grown to high-titer, (iii) the ability to infect a broad range of mammalian cell types, and (iv) the ability to be used with many different promoters including ubiquitous, tissue specific, and regulatable promoters. In addition, adenoviruses can be administered by intravenous injection, because the viruses are stable in the blood-stream.

Using adenovirus vectors where portions of the adenovirus genome are deleted, inserts are incorporated into the viral DNA by direct ligation or by homologous recombination with a co-transfected plasmid. In an exemplary system, the essential E1 gene is deleted from the viral vector, and the virus will not replicate unless the E1 gene is provided by the host cell. When intravenously administered to intact animals, adenovirus primarily targets the liver. Although an adenoviral delivery system with an E1 gene deletion cannot replicate in the host cells, the host's tissue will express and process an encoded heterologous protein. Host cells will also secrete the heterologous protein if the corresponding gene includes a secretory signal sequence. Secreted proteins will enter the circulation from tissue that expresses the heterologous gene (e.g., the highly vascularized liver).

Moreover, adenoviral vectors containing various deletions of viral genes can be used to reduce or eliminate immune responses to the vector. Such adenoviruses are E1-deleted, and in addition, contain deletions of E2A or E4 (Lusky et al., *J. Virol.* 72:2022 (1998); Raper et al., *Human Gene Therapy* 9:671 (1998)). The deletion of E2b has also been reported to reduce immune responses (Amalfitano et al., *J. Virol.* 72:926 (1998)). By deleting the entire adenovirus genome, very large inserts of heterologous DNA can be accommodated. Generation of so called "gutless" adenoviruses, where all viral genes are deleted, are particularly advantageous for insertion of large inserts of heterologous DNA (for a review, see Yeh. and Perricaudet, *FASEB J.* 11:615 (1997)).

High titer stocks of recombinant viruses capable of expressing a therapeutic gene can be obtained from infected mammalian cells using standard methods. For example, recombinant herpes simplex virus can be prepared in Vero cells, as described by Brandt et al., *J. Gen. Virol.* 72:2043 (1991), Herold et al., *J. Gen. Virol.* 75:1211 (1994), Visalli and Brandt, *Virology* 185:419 (1991), Grau et al., *Invest. Ophthalmol. Vis. Sci.* 30:2474 (1989), Brandt et al., *J. Virol. Meth.* 36:209 (1992), and by Brown and MacLean (eds.), *HSV Virus Protocols* (Humana Press 1997).

Alternatively, an expression vector comprising a TACI-immunoglobulin gene can be introduced into a subject's cells by lipofection in vivo using liposomes. Synthetic cationic lipids can be used to prepare liposomes for in vivo transfection of a gene encoding a marker (Feigner et al., *Proc. Nat'l Acad. Sci. USA* 84:7413 (1987); Mackey et al., *Proc. Nat'l Acad. Sci. USA* 85:8027 (1988)). The use of lipofection to introduce exogenous genes into specific organs in vivo has

certain practical advantages. Liposomes can be used to direct transfection to particular cell types, which is particularly advantageous in a tissue with cellular heterogeneity, such as the pancreas, liver, kidney, and brain. Lipids may be chemically coupled to other molecules for the purpose of targeting. Targeted peptides (e.g., hormones or neurotransmitters), proteins such as antibodies, or non-peptide molecules can be coupled to liposomes chemically.

Electroporation is another alternative mode of administration. For example, Aihara and Miyazaki, *Nature Biotechnol*ogy 16:867 (1998), have demonstrated the use of in vivo electroporation for gene transfer into muscle.

In general, the dosage of a composition comprising a therapeutic vector having a TACI-immunoglobulin nucleotide acid sequence, such as a recombinant virus, will vary depending upon such factors as the subject's age, weight, height, sex, general medical condition and previous medical history. Suitable routes of administration of therapeutic vectors include intravenous injection, intraarterial injection, intraperitoneal injection, intramuscular injection, intratumoral injection, and injection into a cavity that contains a tumor. As an illustration, 20 Horton et al., Proc. Nat'l Acad. Sci. USA 96:1553 (1999), demonstrated that intramuscular injection of plasmid DNA encoding interferon-α produces potent antitumor effects on primary and metastatic tumors in a murine model.

A composition comprising viral vectors, non-viral vectors, 25 or a combination of viral and non-viral vectors of the present invention can be formulated according to known methods to prepare pharmaceutically useful compositions, whereby vectors or viruses are combined in a mixture with a pharmaceutically acceptable carrier. As noted above, a composition, 30 such as phosphate-buffered saline is said to be a "pharmaceutically acceptable carrier" if its administration can be tolerated by a recipient subject. Other suitable carriers are wellknown to those in the art (see, for example, Remington's Pharmaceutical Sciences, 19th Ed. (Mack Publishing Co. 35 1995), and Gilman's the Pharmacological Basis of Therapeutics, 7th Ed. (MacMillan Publishing Co. 1985)).

For purposes of therapy, a therapeutic gene expression vector, or a recombinant virus comprising such a vector, and subject in a therapeutically effective amount. A combination of an expression vector (or virus) and a pharmaceutically acceptable carrier is said to be administered in a "therapeutically effective amount" if the amount administered is physiologically significant. An agent is physiologically significant 45 if its presence results in a detectable change in the physiology of a recipient subject. For example, an agent used to treat inflammation is physiologically significant if its presence alleviates the inflammatory response. As another example, an agent used to inhibit the growth of tumor cells is physiologi- 50 cally significant if the administration of the agent results in a decrease in the number of tumor cells, decreased metastasis, a decrease in the size of a solid tumor, or increased necrosis of

When the subject treated with a therapeutic gene expres- 55 sion vector or a recombinant virus is a human, then the therapy is preferably somatic cell gene therapy. That is, the preferred treatment of a human with a therapeutic gene expression vector or a recombinant virus does not entail introducing into cells a nucleic acid molecule that can form part of 60 a human germ line and be passed onto successive generations (i.e., human germ line gene therapy).

10. Production of Transgenic Mice

Transgenic mice can be engineered to over-express nucleic acid sequences encoding TACI-immunoglobulin fusion pro42

teins in all tissues, or under the control of a tissue-specific or tissue-preferred regulatory element. These over-producers of TACI-immunoglobulin fusion proteins can be used to characterize the phenotype that results from over-expression, and the transgenic animals can serve as models for human disease caused by excess TACI receptor protein. Transgenic mice that over-express TACI-immunoglobulin fusion proteins also provide model bioreactors for production of TACI-immunoglobulin fusion proteins in the milk or blood of larger animals. Methods for producing transgenic mice are well-known to those of skill in the art (see, for example, Jacob, "Expression and Knockout of Interferons in Transgenic Mice," in Overexpression and Knockout of Cytokines in Transgenic Mice, Jacob (ed.), pages 111-124 (Academic Press, Ltd. 1994), Monastersky and Robl (eds.), Strategies in Transgenic Animal Science (ASM Press 1995), and Abbud and Nilson, "Recombinant Protein Expression in Transgenic Mice," in Gene Expression Systems: Using Nature for the Art of Expression, Fernandez and Hoeffler (eds.), pages 367-397 (Academic Press, Inc. 1999)).

For example, a method for producing a transgenic mouse that expresses a nucleic acid sequence that encodes a TACIimmunoglobulin fusion protein can begin with adult, fertile males (studs) (B6C3f1, 2 to 8 months of age (Taconic Farms, Germantown, N.Y.)), vasectomized males (duds) (B6D2f1, 2 to 8 months, (Taconic Farms)), prepubescent fertile females (donors) (B6C3f1, 4 to 5 weeks, (Taconic Farms)) and adult fertile females (recipients) (B6D2f1, 2 to 4 months, (Taconic Farms)). The donors are acclimated for one week and then injected with approximately 8 IU/mouse of Pregnant Mare's Serum gonadotrophin (Sigma Chemical Company; St. Louis, Mo.) I.P., and 46-47 hours later, 8 IU/mouse of human Chorionic Gonadotropin (hCG (Sigma)) I.P. to induce superovulation. Donors are mated with studs subsequent to hormone injections. Ovulation generally occurs within 13 hours of hCG injection. Copulation is confirmed by the presence of a vaginal plug the morning following mating.

Fertilized eggs are collected under a surgical scope. The a pharmaceutically acceptable carrier are administered to a 40 oviducts are collected and eggs are released into urinanalysis slides containing hyaluronidase (Sigma). Eggs are washed once in hyaluronidase, and twice in Whitten's W640 medium (described, for example, by Menino and O'Claray, Biol. Reprod. 77:159 (1986), and Dienhart and Downs, Zygote 4:129 (1996)) that has been incubated with 5% CO₂, 5% O₂, and 90% N_2 at 37° C. The eggs are then stored in a 37° C./5% CO₂ incubator until microinjection.

> Ten to twenty micrograms of plasmid DNA containing a TACI-immunoglobulin fusion protein encoding sequence is linearized, gel-purified, and resuspended in 10 mM Tris-HCl (pH 7.4), 0.25 mM EDTA (pH 8.0), at a final concentration of 5-10 nanograms per microliter for microinjection. For example, the TACI-immunoglobulin fusion protein encoding sequences can encode a TACI polypeptide with deletion of amino acid residues 1 to 29 and 111 to 154 of SEQ ID NO:2, and an Fc5 immunoglobulin moiety.

> Plasmid DNA is microinjected into harvested eggs contained in a drop of W640 medium overlaid by warm, CO₂equilibrated mineral oil. The DNA is drawn into an injection needle (pulled from a 0.75 mm ID, 1 mm OD borosilicate glass capillary), and injected into individual eggs. Each egg is penetrated with the injection needle, into one or both of the haploid pronuclei.

Picoliters of DNA are injected into the pronuclei, and the injection needle withdrawn without coming into contact with the nucleoli. The procedure is repeated until all the eggs are injected. Successfully microinjected eggs are transferred into

an organ tissue-culture dish with pre-gassed W640 medium for storage overnight in a 37° C./5% CO₂ incubator.

The following day, two-cell embryos are transferred into pseudopregnant recipients. The recipients are identified by the presence of copulation plugs, after copulating with vasectomized duds. Recipients are anesthetized and shaved on the dorsal left side and transferred to a surgical microscope. A small incision is made in the skin and through the muscle wall in the middle of the abdominal area outlined by the ribcage, the saddle, and the hind leg, midway between knee and spleen. The reproductive organs are exteriorized onto a small surgical drape. The fat pad is stretched out over the surgical drape, and a baby serrefine (Roboz, Rockville, Md.) is attached to the fat pad and left hanging over the back of the 15 mouse, preventing the organs from sliding back in.

With a fine transfer pipette containing mineral oil followed by alternating W640 and air bubbles, 12-17 healthy two-cell embryos from the previous day's injection are transferred into the recipient. The swollen ampulla is located and holding the 20 oviduct between the ampulla and the bursa, a nick in the oviduct is made with a 28 g needle close to the bursa, making sure not to tear the ampulla or the bursa.

The pipette is transferred into the nick in the oviduct, and the embryos are blown in, allowing the first air bubble to 25 escape the pipette. The fat pad is gently pushed into the peritoneum, and the reproductive organs allowed to slide in. The peritoneal wall is closed with one suture and the skin closed with a wound clip. The mice recuperate on a 37° C. slide warmer for a minimum of four hours.

The recipients are returned to cages in pairs, and allowed 19-21 days gestation. After birth, 19-21 days postpartum is allowed before weaning. The weanlings are sexed and placed into separate sex cages, and a 0.5 cm biopsy (used for genotyping) is snipped off the tail with clean scissors.

Genomic DNA is prepared from the tail snips using, for example, a QIAGEN DNEASY kit following the manufacturer's instructions. Genomic DNA is analyzed by PCR using primers designed to amplify a nucleic acid sequence encoding a TACI-immunoglobulin fusion protein or a selectable 40 marker gene that was introduced in the same plasmid. After animals are confirmed to be transgenic, they are back-crossed into an inbred strain by placing a transgenic female with a wild-type male, or a transgenic male with one or two wildtype female(s). As pups are born and weaned, the sexes are 45 separated, and their tails snipped for genotyping.

To check for expression of a transgene in a live animal, a partial hepatectomy is performed. A surgical prep is made of the upper abdomen directly below the zyphoid process. Using sterile technique, a small 1.5-2 cm incision is made below the 50 human IgG1 (the hinge region and the CH₂ and CH₃ domains) sternum and the left lateral lobe of the liver exteriorized. Using 4-0 silk, a tie is made around the lower lobe securing it outside the body cavity. An atraumatic clamp is used to hold the tie while a second loop of absorbable Dexon (American Cyanamid; Wayne, N.J.) is placed proximal to the first tie. A 55 distal cut is made from the Dexon tie and approximately 100 mg of the excised liver tissue is placed in a sterile petri dish. The excised liver section is transferred to a 14 ml polypropylene round bottom tube and snap frozen in liquid nitrogen and then stored on dry ice. The surgical site is closed with suture 60 and wound clips, and the animal's cage placed on a 37° C. heating pad for 24 hours post operatively. The animal is checked daily post operatively and the wound clips removed 7-10 days after surgery. The expression level of TACI-immunoglobulin fusion protein mRNA is examined for each transgenic mouse using an RNA solution hybridization assay or polymerase chain reaction.

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Using the general approach described above, transgenic mice have been produced that express significant levels of TACI-immunoglobulin fusion protein in milk. In this particular case, the TACI-immunoglobulin fusion protein encoding sequence encoded a TACI polypeptide with deletion of amino acid residues 1 to 29 and 111 to 154 of SEQ ID NO:2, and an Fc5 immunoglobulin moiety.

The present invention, thus generally described, will be understood more readily by reference to the following examples, which are provided by way of illustration and is not intended to be limiting of the present invention.

EXPERIMENTAL

Example 1

Construction of Nucleic Acid Molecules that Encode TACI-Fc Proteins

Nucleic acid molecules encoding human TACI were obtained during the expression cloning of the receptors for ZTNF4 as described by Gross et al., Nature 404:995 (2000). The coding sequences contained in the TACI-Fc expression constructs were generated by overlap PCR, using standard techniques (see, for example, Horton et al., Gene 77:61 (1989)). Human TACI cDNA and Fc cDNA were used as starting templates for the PCR amplifications. PCR primers were designed to yield the desired coding sequence 5' and 3' ends and to introduce restriction enzyme recognition sites to facilitate insertion of these coding sequences into the expression vectors. The TACI-Fc coding sequences were inserted into expression vectors that included a functional murine dihydrofolate reductase gene. One expression vector also contained a cytomegalovirus promoter to direct the expression of the recombinant protein transgene, an immunoglobulin intron, a tissue plasminogen activator signal sequence, an internal ribosome entry sequence, a deleted CD8 cistron for surface selection of transfected cells, and yeast expression elements for growth of the plasmid in yeast cells.

One approach that was used to produce TACI-Fc fusion proteins is illustrated by the method used to construct TACI-Fc4. Other TACI-Fc fusion proteins were produced by inserting nucleotide sequences that encode a TACI-Fc fusion protein into a mammalian expression vector, and introducing that expression vector into mammalian cells.

A. Ig γ1 Fc4 Fragment Construction

To prepare the TACI-Fc4 fusion protein, the Fc region of was modified to remove Fcy1 receptor (FcyRI) and complement (C1q) binding functions. This modified version of human IgG1 Fc was designated "Fc4."

The Fc region was isolated from a human fetal liver library (Clontech) PCR using oligo primers 5' ATCAGCGGAA TTCAGATCTT CAGACAAAAC TCACACATGC CCAC 3' (SEQ ID NO:7) and 5' GGCAGTCTCT AGATCATTTA CCCGGAGACA GGGAG 3' (SEQ ID NO:8). Mutations within the Fc region were introduced by PCR to reduce FcyRI binding. The FcγRI binding site (Leu-Leu-Gly-Gly; amino acid residues 38 to 41 of SEQ ID NO:6, which correspond to EU index positions 234 to 237) was mutated to Ala-Glu-Gly-Ala to reduce FcyR1 binding (see, for example, Duncan et al., Nature 332:563 (1988); Baum et al., EMBO J. 13:3992 (1994)). Oligonucleotide primers 5' CCGTGCCCAG CAC-CTGAAGC CGAGGGGCA CCGTCAGTCT TCCTCT-TCCC C 3' (SEQ ID NO:9) and 5' GGATTCTAGA TTATT-

TACCC GGAGACAGGGA 3' (SEQ ID NO:10) were used to introduce the mutation. To a 50 μl final volume was added 570 ng of IgFc template, 5 μl of 10× Pfu reaction Buffer (Stratagene), 8 μl of 1.25 mM dNTPs, 31 μl of distilled water, 2 μl of 20 mM oligonucleotide primers. An equal volume of mineral oil was added and the reaction was heated to 94° C. for 1 minute. Pfu polymerase (2.5 units, Stratagene) was added followed by 25 cycles at 94° C. for 30 seconds, 55° C. for 30 seconds, 72° C. for 1 minute followed by a 7 minute extension at 72° C. The reaction products were fractioned by electrophoresis, and the band corresponding to the predicted size of about 676 base pairs was detected. This band was excised from the gel and recovered using a QIAGEN QIAquickTM Gel Extraction Kit (Qiagen) according to the manufacturer's instructions.

PCR was also used to introduce a mutation of Ala to Ser (amino acid residue 134 of SEQ ID NO:6, which corresponds to EU index position 330) and Pro to Ser (amino acid residue 135 of SEQ ID NO:6, which corresponds to EU index position 331) to reduce complement C1q binding or complement 20 fixation (Duncan and Winter, Nature 332:788 (1988)). Two first round reactions were performed using the FcyRI binding side-mutated IgFc sequence as a template. To a 5 μ1 final volume was added 1 μl of FcγRI binding site mutated IgFc template, 5 µl of 10×Pfu Reaction Buffer (Stratagene), 8 µl of 25 1.25 mM dNTPs, 31 of µl distilled water, 2 µl of 20 mM 5' GGTGGCGGCT CCCAGATGGG TCCTGTCCGA GCCCAGATCT TCAGACAAAA CTCAC 3' (SEQ ID NO:11), a 5' primer beginning at nucleotide 36 of SEQ ID NO:5, and 2 µl of 20 mM 5' TGGGAGGGCT TTGTTGGA 3' (SEQ ID NO:12), a 3' primer beginning at the complement of nucleotide 405 of SEQ ID NO:5. The second reaction contained 2 µl each of 20 mM stocks of oligonucleotide primers TCCAACAAG CCCTCCCATC CTCCATCGAG AAAACCATCT CC 3' (SEQ ID NO:13), a 5' primer begin- 35 ning at nucleotide 388 of SEQ ID NO:5 and 5' GGATG-GATCC ATGAAGCACC TGTGGTTCTT CCTCCTGCTG GTGGCGGCTC CCAGATG 3' (SEQ ID NO:14), a 3' primer, to introduce the Ala to Ser mutation, XbaI restriction site and stop codon. An equal volume of mineral oil was added and the 40 reactions were heated to 94° C. for 1 minute. Pfu polymerase (2.5 units, Stratagene) was added followed by 25 cycles at 94° C. for 30 seconds, 55° C. for 30 seconds, 72° C. for 2 minutes followed by a 7 minute extension at 72° C. The reaction products were fractionated by electrophoresis, and bands cor- 45 responding to the predicted sizes, about 370 and about 395 base pairs respectively, were detected. The bands were excised from the gel and extracted using a QIAGEN QIAquick™ Gel Extraction Kit (Qiagen) according to the manufacturer's instructions.

A second round reaction was performed to join the above fragments and add the 5' BamHI restriction site and a signal sequence from the human immunoglobulin JBL 2'C₁ heavy chain variable region (Cogne et al., Eur. J. Immunol. 18:1485 (1988)). To a 50 μ l final volume was added 3 μ l of distilled 55 water, 8 μl of 1.25 mM dNTPs, 5 μl of 10×Pfu polymerase reaction buffer (Stratagene) and 1 µl each of the two first two PCR products. An equal volume of mineral oil was added and the reaction was heated to 94° C. for 1 minute. Pfu polymerase (2.5 units, Stratagene) was added followed by 5 cycles 60 at 94° C. for 30 seconds, 55° C. for 30 seconds, and 72° C. for 2 minutes. The temperature was again brought to 94° C. and 2 µl each of 20 mM stocks of 5' GGATGGATCC ATGAAG-CACC TGTGGTTCTT CCTCCTGCTG GTGGCGGCTC CCAGATG 3' (SEQ ID NO:14), a 5' primer beginning at 65 nucleotide 1 of SEQ ID NO:5, and 5' GGATTCTAGA TTATTTACCC GGAGACAGGGA 3' (SEQ ID NO:10) were

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added followed by 25 cycles at 94° C. for 30 seconds, 55° C. for 30 seconds and 72° C. for 2 minutes, and a final 7 minute extension at 72° C. A portion of the reaction was visualized using gel electrophoresis. A 789 base pair band corresponding the predicted size was detected.

B. TACI-Fc4 Expression Vector Construction

Expression plasmids comprising a coding region for TACIat 72° C. The reaction products were fractioned by electro- 10 Fc4 fusion protein were constructed via homologous recombination in yeast. A fragment of TACI cDNA was isolated using PCR that included the polynucleotide sequence from nucleotide 14 to nucleotide 475 of SEQ ID NO:1. The two primers used in the production of the TACI fragment were: (1) a primer containing 40 base pairs of the 5' vector flanking sequence and 17 base pairs corresponding to the amino terminus of the TACI fragment (5' CTCAGCCAGG AAATC-CATGC CGAGTTGAGA CGCTTCCGTA GAGTGG CCTGGGCCG 3'; SEQ ID NO:15); (2) 40 base pairs of the 3' end corresponding to the flanking Fc4 sequence and 17 base pairs corresponding to the carboxyl terminus of the TACI fragment (5' GCATGTGTGA GTTTTGTCTG AAGATCTGGG CTCCTTCAGC CCCGGGAG 3'; SEQ ID NO:16). To a 100 µl final volume was added 10 ng of TACI template, 10 µl of 10× Taq polymerase Reaction Buffer (Perkin Elmer), 8 μl of 2.5 nM dNTPs, 78 μl of distilled water, 2 μl each of 20 mM stocks of the oligonucleotide primers, and Taq polymerase (2.5 units, Life Technology). An equal volume of mineral oil was added and the reaction was heated to 94° C. for 2 minutes, followed by 25 cycles at 94° C. for 30 seconds, 65° C. for 30 seconds, 65° C. for 30 seconds, 72° C. for 1 minute followed by a 5 minute extension at 72° C.

The fragment containing the cDNA encoding the Fc4 fragment was constructed in a similar manner. The two primers used in the production of the Fc4 fragment were (upstream and downstream), an oligonucleotide primer containing 40 base pairs of the 5' TACI flanking sequence and 17 base pairs corresponding to the amino terminus of the Fc4 fragment (5' GCACAGAGGC TCAGAAGCAA GTCCAGCTCT CCCGGGGCTG AAGGAGCCCA GATCTTCAGA 3'; SEQ ID NO:17); and an oligonucleotide primer containing 40 base pairs of the 3' end corresponding to the flanking vector sequence and 17 base pairs corresponding to the carboxyl terminus of the Fc4 fragment (5' GGGGTGGGTA CAAC-CCCAGA GCTGTTTTAA TCTAGATTAT TTACCCGGAG ACAGGG 3'; SEQ ID NO:18). To a 100 µl final volume was added 10 ng of Fc4 template described above, 10 µl 10× Taq polymerase Reaction Buffer (Perkin Elmer), 8 µl of 2.5 nM dNTPs, 78 μl of distilled water, 2 μl each of 20 mM stocks of the oligonucleotides, and Taq polymerase (2.5 units, Life Technology). An equal volume of mineral oil was added and the reaction was heated to 94° C. for 2 minutes, then 25 cycles at 94° C. for 30 seconds, 65° C. for 30 seconds, 72° C. for 1 minute followed by a 5 minute extension at 72° C.

Ten microliters of each of the 100 µl PCR reactions described above were run on a 0.8% LMP agarose gel (Seaplaque GTG) with 1×TBE buffer for analysis. The remaining 90 µl of each PCR reaction was precipitated with the addition of 5 µl of 1 M sodium chloride and 250 µl of absolute ethanol. The plasmid pZMP6 was cleaved with Sinai to linearize it at the polylinker. Plasmid pZMP6 was derived from the plasmid pCZR199 (American Type Culture Collection, Manassas, Va., ATCC#98668) and is a mammalian expression vector containing an expression cassette having the cytomegalovirus immediate early promoter, a consensus intron from the variable region of mouse immunoglobulin heavy chain locus, multiple restriction sites for insertion of coding sequences, a

stop codon and a human growth hormone terminator. The plasmid also has an *E. coli* origin of replication, a mammalian selectable marker expression unit having an SV40 promoter, enhancer and origin of replication, a dihydrofolate reductase gene and the SV40 terminator. The vector pZMP6 was constructed from pCZR199 by replacement of the metallothionein promoter with the cytomegalovirus immediate early promoter, and the Kozac sequences at the 5' end of the open reading frame.

One hundred microliters of competent yeast cells ($S.\ cervevisiae$) were combined with $10\,\mu l$ containing approximately $1\,\mu g$ of the TACI extracellular domain and the Fc4 PCR fragments, and $100\,n g$ of SmaI digested pZMP6 vector and transferred to a $0.2\,cm$ electroporation cuvette. The yeast/DNA mixtures were electropulsed at $0.75\,kV$ ($5\,kV/cm$), ∞ 15 ohms, $25\,\mu F$. To each cuvette was added $600\,\mu l$ of $1.2\,M$ sorbitol and the yeast were plated in two $300\,\mu l$ aliquots onto to URA-D plates and incubated at $30^{\circ}\,C$.

After about 48 hours, the Ura+ yeast transformants from a single plate were resuspended in 1 ml of water and spun 20 briefly to pellet the yeast cells. The cell pellet was resuspended in 1 ml of lysis buffer (2% Triton X-100, 1% SDS, 100 mM NaCl, 10 mM Tris, pH 8.0, 1 mM EDTA). Five hundred microliters of the lysis mixture was added to an Eppendorf tube containing 300 μl acid washed glass beads and 200 μl 25 phenol-chloroform, vortexed for 1 minute intervals two or three times, followed by a 5 minute spin in a Eppendorf centrifuge at maximum speed. Three hundred microliters of the aqueous phase was transferred to a fresh tube, and the DNA precipitated with 600 μl of ethanol, followed by centrifugation for 10 minutes at 4° C. The DNA pellet was resuspended in 100 μl of water.

Transformation of electrocompetent *E. coli* cells (DH10B, GibcoBRL) was performed with 0.5-2 ml yeast DNA prep and 40 µl of DH10B cells. The cells were electropulsed at 2.0 ³⁵ kV, 25 mF and 400 ohms. Following electroporation, 1 ml of SOC (2% Bacto-Tryptone (Difco, Detroit, Mich.), 0.5% yeast extract (Difco), 10 mM NaCl, 2.5 mM KCl, 10 mM MgCl₂, 10 mM MgSO₄, 20 mM glucose) were plated in 250 µl aliquots on four LB AMP plates (LB broth (Lennox), 1.8% ⁴⁰ Bacto-Agar (Difco), 100 mg/L Ampicillin).

Individual clones harboring the correct expression construct for TACI-Fc4 were identified by restriction digest to verify the presence of the insert and to confirm that the various DNA sequences have been joined correctly to one another. 45 The insert of positive clones were subjected to sequence analysis. Larger scale plasmid DNA is isolated using the Qiagen Maxi kit (Qiagen) according to manufacturer's instructions.

C. Construction of Fc5, Fc6, and Fc7

In Fc5, the Arg residue at EU index position 218 was changed back to a Lys residue. Wild-type human Ig γ1 con-

tains a lysine at this position. Briefly, nucleic acid molecules 55 encoding Fc5 were produced using oligonucleotide primers 5' GAGCCCAAATCTTCAGACAAAACTCACACATGCCCA 3' (SEQ ID NO:19) and 5' TAATTGGCGCGCCTCTAGATTATTTACCCGGAGACA 3' (SEQ ID NO:20). The conditions of the PCR amplification were as follows. To 60 a 50 μ l final volume was added 236 ng of Fc4 template, 5 μ l of 10 Pfu reaction Buffer (Stratagene), 4 μ l of 2.5 mM dNTPs, 1 μ l of 20 μ M of each of the oligonucleotides, and 1 μ l of Pfu polymerase (2.5 units, Stratagene). The amplification thermal profile consisted of 94° C. for 2 minutes, 5 cycles at 94° C. for 515 seconds, 42° C. for 20 seconds, 72° C. for 45 seconds, 20 cycles at 94° C. for 15 seconds, 72° C. for 1 minute 20

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seconds, followed by a 7 minute extension at 72° C. The reaction product was fractionated by agarose gel electrophoresis, and the band corresponding to the predicted size of about 718 base pairs was detected. The band was excised from the gel and recovered using a QIAGEN QIAquick Gel Extraction Kit (Qiagen) according to the manufacturer's instructions.

Fc6 is identical to Fc5 except that the carboxyl terminal lysine codon has been eliminated. As in Fc4 and Fc5 above, the stop codon in the Fc6 sequence was changed to TAA. Fc6 was generated from template DNA that encoded Fc5 using oligonucleotide primers 5' GAGCCCAAAT CTTCAGA-CAA AACTCACACA TGCCCA 3' (SEQ ID NO:19) and 5' GGCGCGCCTC TAGATTAACC CGGAGACAGG GAGAGGC 3' (SEQ ID NO:21).

Fc7 is identical to the wild-type γ1 Fc except for an amino acid substitution at EU index position Asn 297 located in the $C_H 2$ domain. As 297 was mutated to a Gln residue to prevent the attachment of N-linked carbohydrate at that residue position. As above, the stop codon in the Fc7 sequence was changed to TAA. Fc7 was generated by overlap PCR using a wild-type human IgGγ1 Fc cDNA as the template and oligonucleotide primers 5' GAGCCCAAATCTTGCGA-CAAAACTCACA 3' (SEQ ID NO:22) and 5' GTACGT-GCTTTGGTACTGCTCCTCCCGCGGCTT 3' (SEQ ID NO:23) to generate the 5' half of Fc7, and oligonucleotide primers 5' CAGTACCAAAGCACGTACCGTGTGGTCA 3' (SEQ ID NO:24) and 5' TAATTGGCGCGCCTCTAGAT-TATTTACCCGGAGACA 3' (SEQ ID NO:20) to generate the 3' half of Fc7. The two PCR products were combined and amplified using oligonucleotide primers 5' GAGC-CCAAATCTTGCGACAAAACTCACA 3' (SEQ ID NO:22) and TAATTGGCGCGCCTCTAGATTATTTAC-CCGGAGACA 3' (SEQ ID NO:20).

All the resultant PCR products were gel purified, cloned, and verified by DNA sequence analysis.

D. Construction of Amino-Truncated TACI-Fc Fusion Proteins

Four amino terminal truncated versions of TACI-Fc were generated. All four had a modified human tissue plasminogen activator signal sequence (SEQ ID NO:25) fused to amino acid residue number 30 of SEQ ID NO:2. However, the four proteins differed in the location of point in which the Fc5 was fused to the TACI amino acid sequence of SEQ ID NO:2.

Table 3 outlines the structures of the four fusion proteins.

TABLE 3

	TACI Fusion Proteins									
5	Designation of TACI-Fc	TACI amino acid residues								
	TACI(d1-29)-Fc5 TACI(d1-29, d107-154)-Fc5 TACI(d1-29, d111-154)-Fc5 TACI(d1-29, d120-154)-Fc5	30 to 154 of SEQ ID NO: 2 30 to 106 of SEQ ID NO: 2 30 to 110 of SEQ ID NO: 2 30 to 119 of SEQ ID NO: 2								

Protein encoding expression cassettes were generated by overlap PCR using standard techniques (see, for example, Horton et al., *Gene* 77:61 (1989)). A nucleic acid molecule encoding TACI and a nucleic acid molecule encoding Fc5 were used as PCR templates. Oligonucleotide primers are identified in Tables 4 and 5.

20

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TABLE 4

Oligonucleotide Primers Used to Produce TACI Fusion Proteins						
	Oligonucleotide Designations					
Designation of TACI-Fc	5' TACI	3' TACI	5' Fc5	3' Fc5		
TACI(d1-29)-Fc5 TACI(d1-29, d107-154)- Fc5	ZC24,903 ZC24,903	ZC24,955 ZC24,951	ZC24,952 ZC24,949	ZC24,946 ZC24,946		
TACI(d1-29, d111-154)- Fc5	ZC24,903	ZC28,978	ZC28,979	ZC24,946		
TACI(d1-29, d120-154)- Fc5	ZC24,903	ZC28,981	ZC28,980	ZC24,946		

TABLE 5

	Oliqonucleotide Sequences	
Primer	Nucleotide Sequence	SEQ ID NO.
ZC24,903	5' TATTAGGCCGGCCACCATGGATGCAATGA 3'	40
ZC24,955	5' TGAAGATTTGGGCTCCTTGAGACCTGGGA 3'	41
ZC24,952	5' TCCCAGGTCTCAAGGAGCCCAAATCTTCA 3'	42
ZC24,946	5' TAATTGGCGCGCCTCTAGATTATTTACCCGGA GACA 3'	20
ZC24,951	5' TGAAGATTTGGGCTCGTTCTCACAGAAGTA 3'	43
ZC24,949	5' ATACTTCTGTGAGAACGAGCCCAAATCT TCA 3'	44
ZC28,978	5' TTTGGGCTCGCTCCTGAGCTTGTTCTCACA 3'	45
ZC28,979	5' CTCAGGAGCGAGCCCAAATCTTCAGACA 3'	46
ZC28,981	5' TTTGGGCTCCCTGAGCTCTGGTGGAA 3'	47
ZC28,980	5' GAGCTCAGGGAGCCCAAATCTTCAGACA 3'	48

The first round of PCR amplifications consisted of two reactions for each of the four amino terminal truncated versions. The two reactions were performed separately using the 5' and 3' TACI oligonucleotides in one reaction, and the 5' and The conditions of the first round PCR amplification were as follows. To a 25 µl final volume was added approximately 200 ng template DNA, 2.5 μl 10×Pfu reaction Buffer (Stratagene), $2 \mu l$ of $2.5 \, mM \, dNTPs$, $0.5 \, \mu l$ of $20 \, \mu M$ each 5' oligonucleotide and 3' oligonucleotide, and 0.5 µl Pfu polymerase (2.5 units, 50 Stratagene). The amplification thermal profile consisted of 94° C. for 3 minutes, 35 cycles at 94° C. for 15 seconds, 50° C. for 15 seconds, 72° C. for 2 minutes, followed by a 2 minute extension at 72° C. The reaction products were fractionated by agarose gel electrophoresis, and the bands corre- 55 Biosciences, Lenexa, Kans.), 4 mM L-Glutamine (JRH Biosponding to the predicted sizes were excised from the gel and recovered using a QIAGEN QIAQUICK Gel Extraction Kit (Qiagen), according to the manufacturer's instructions.

The second round of PCR amplification, or overlap PCR amplification reaction, was performed using the gel purified 60 fragments from the first round PCR as DNA template. The conditions of the second round PCR amplification were as follows. To a 25 µl final volume was added approximately 10 ng template DNA each of the TACI fragment and the Fc5 fragment, 2.5 µl 10×Pfu reaction Buffer (Stratagene), 2 µl of 65 2.5 mM dNTPs, 0.5 µl of 20 µM each ZC24,903 (SEQ ID NO:40) and ZC24,946 (SEQ ID NO:20) and 0.5 µl Pfu poly-

merase (2.5 units, Stratagene). The amplification thermal profile consisted of 94° C. for 1 minute, 35 cycles at 94° C. for 15 seconds, 55° C. for 15 seconds, 72° C. for 2 minutes, followed by a 2 minute extension at 72° C. The reaction products were fractionated by agarose gel electrophoresis, and the bands corresponding to the predicted sizes were excised from the gel and recovered using a OIAGEN OIAOUICK Gel Extraction Kit (Qiagen), according to the manufacturer's instruc-

Each of the four versions of the amino terminal truncated TACI-Fc PCR products were separately cloned using Invitrogen's ZEROBLUNT TOPO PCR Cloning Kit following the manufacturer's recommended protocol. Table 6 identifies the nucleotide and amino acid sequences of these TACI-Fc con-

TABLE 6

	SEQ	ID Nos.
Designation of TACI-Fc	Nucleotide	Amino Acid
TACI(d1-29)-Fc5	49	50
TACI(d1-29, d107-154)-Fc5	51	52
TACI(d1-29, d111-154)-Fc5	53	54
TACI(d1-29, d120-154)-Fc5	55	56

After the nucleotide sequences were verified, plasmids 30 comprising each of the four versions of the amino terminal truncated TACI-Fc fusions were digested with FseI and AscI to release the amino acid encoding segments. The FseI-AscI fragments were ligated into a mammalian expression vector containing a CMV promoter and an SV40 poly A segment. Expression vectors were introduced into Chinese hamster ovary cells as described below.

Example 2

Production of TACI-Fc Proteins by Chinese Hamster Ovary Cells

The TACI-Fc expression constructs were used to transfect, via electroporation, suspension-adapted Chinese hamster 3' Fc5 oligonucleotides in another reaction for each version. 45 ovary (CHO) DG44 cells grown in animal protein-free medium (Urlaub et al., Som. Cell. Molec. Genet. 12:555 (1986)). CHO DG44 cells lack a functional dihydrofolate reductase gene due to deletions at both dihydrofolate reductase chromosomal locations. Growth of the cells in the presence of increased concentrations of methotrexate results in the amplification of the dihydrofolate reductase gene, and the linked recombinant protein-encoded gene on the expression

> CHO DG44 cells were passaged in PFCHO media (JRH sciences), and 1× hypothanxine-thymidine supplement (Life Technologies), and the cells were incubated at 37° C. and 5% CO₂ in Corning shake flasks at 120 RPM on a rotating shaker platform. The cells were transfected separately with linearized expression plasmids. To ensure sterility, a single ethanol precipitation step was performed on ice for 25 minutes by combining 200 µg of plasmid DNA in an Eppendorf tube with 20 µl of sheared salmon sperm carrier DNA (5'→3' Inc. Boulder, Colo., $10\,mg/ml),\,22\,\mu l$ of 3M NaOAc (pH 5.2), and 484 μl of 100% ethanol (Gold Shield Chemical Co., Hayward, Calif.). After incubation, the tube was centrifuged at 14,000 RPM in a microfuge placed in a 4° C. cold room, the super-

natant removed and the pellet washed twice with 0.5 ml of 70% ethanol and allowed to air dry.

The CHO DG44 cells were prepared while the DNA pellet was drying by centrifuging 10⁶ total cells (16.5 ml) in a 25 ml conical centrifuge tube at 900 RPM for 5 minutes. The CHO ⁵ DG44 cells were resuspended into a total volume of 300 µl of PFCHO growth media, and placed in a Gene-Pulser Cuvette with a 0.4 cm electrode gap (BioRad). The DNA, after approximately 50 minutes of drying time, was resuspended into 500 µl of PFCHO growth media and added to the cells in ¹⁰ the cuvette so that the total volume did not exceed 800 µl and was allowed to sit at room temperature for 5 minutes to decrease bubble formation. The cuvette was placed in a Bio-Rad Gene Pulser II unit set at 0.296 kV (kilovolts) and 0.950 HC (high capacitance) and electroporated immediately.

The cells were incubated 5 minutes at room temperature before placement in 20 ml total volume of PFCHO media in a CoStar T-75 flask. The flask was placed at 37° C. and 5% CO₂ for 48 hours when the cells were then counted by hemocytometer utilizing trypan blue exclusion and put into ²⁰ PFCHO selection media without hypothanxine-thymidine supplement and containing 200 mM methotrexate (Cal Biochem).

Upon recovery of the methotrexate selection process, the conditioned media containing the secreted TACI-Fc proteins ²⁵ were examined by Western Blot analysis.

Example 3

Structural Analysis of TACI-Fc Proteins

In certain cases, TACI-Fc fusion proteins were partially purified before analysis. Conditioned medium from Chinese hamster ovary cultures was sterile-filtered through a 0.22 µm filter and the TACI-Fc protein was captured on a protein A column. The protein A-bound material was eluted and passed over an S-200 size exclusion column for final purification.

Western blot analysis was performed on both conditioned cell medium and purified protein to assess the structural stability of the TACI-Fc proteins. Briefly, protein or supernatant samples were transferred to nitrocellulose membranes and the TACI-Fc proteins were detected using peroxidase conjugated goat anti-mouse IgG2a (Boehringer Mannheim), or peroxidase conjugated goat anti-human IgG Fc specific antisera (Pierce).

Amino terminal amino acid sequence analyses were performed on Models 476A and 494 Protein Sequencer Systems from Perkin Elmer Applied Biosystems Division (Foster City, Calif.). Data analysis was performed with Applied Biosystems Model 610A Data Analysis System for Protein Sequencing, version 2.1a (Applied Biosystems, Inc.). Most supplies and reagents used were from Applied Biosystems, Inc.

Example 4

Functional Analysis of TACI-Fc Proteins

Two approaches were used to examine the binding characteristics of four TACI-Fc proteins with ZTNF4. One approach measured the ability of the TACI-Fc constructs to compete 60 with TACI-coated plates for binding of ¹²⁵I-labeled ZTNF4. In the second approach, increasing concentrations of ¹²⁵I labeled ZTNF4 were incubated with each of the TACI-Fc constructs, and the radioactivity associated with precipitated ZTNF4-TACI-Fc complexes was determined. The TACI-Fc 65 fusion proteins had TACI moieties that lacked the first 29 amino acid residues of the amino acid sequence of SEQ ID

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NO:2. One of the fusion proteins had a TACI moiety with an intact stalk region (TACI (d1-29)-Fc5), whereas three of the TACI-Fc fusion proteins had TACI moieties with various deletions in the stalk region (TACI (d1-29, d107-154)-Fc5; TACI (d1-29, d111-154)-Fc5; TACI (d1-29, d120-154)-Fc5). A. Competitive Binding Assav

ZTNF4 was radiodinated with Iodobeads (Pierce), following standard methods. Briefly, 50 μg of the ZTNF4 was iodinated with 4 mCi of ¹²⁵I using a single Iodobead. The reaction was quenched with a 0.25% solution of bovine serum albumin, and the free ¹²⁵I was removed by gel filtration using a PD-10 column (Pierce). The specific radioactivity of ¹²⁵I-ZTNF4 preparations was determined by trichloroacetic acid precipitation before and after the desalting step.

An N-terminal fragment of the TACI receptor, designated as "TACI-N," was added to 96-well plates (100 μl at 0.1 μg/ml), and incubated overnight at 4° C. The plates were washed, blocked with Superblock (Pierce), and washed again. The TACI-Fc constructs, at various concentrations ranging from 0 to 11.5 ng/ml, were mixed with a fixed concentration of ¹²⁵I-ZTNF4 (20 ng/ml), and incubated for 2 hours at 37° C. on the plate coated with TACI-N. Controls contained either TACI-N in solution, or lacked TACI-Fc. After incubation, the plates were washed and counted. Each determination was performed in triplicate.

The results showed that all TACI-Fc constructs inhibited $^{125}\text{I-ZTNF4}$ binding completely at concentrations of about 100 ng/ml or greater. The TACI-Fc proteins, TACI (d1-29)-Fc5, TACI (d1-29, d111-154)-Fc5, and TACI (d1-29, d120-154)-Fc5, were more effective inhibitors than the TACI-Fc construct, TACI (d1-29, d107-154)-Fc5. An Fc fragment alone did not inhibit binding. IC $_{50}$ values were calculated for each construct in three experiments. The average values for the constructs were: TACI (d1-29)-Fc5: 2.07 nM; TACI (d1-29, d107-154)-Fc5: 4.95 nM; TACI (d1-29, d111-154)-Fc5: 2.31 nM; and TACI (d1-29, d120-154)-Fc5: 2.21 nM. B. Solution Binding Assay

At a concentration of 0.05 nM, each TACI-Fc construct was incubated with 0.4 pM to 1.5 nM ¹²⁵I-ZTNF4 for 30 minutes at room temperature in a total volume of 0.25 ml/tube. A Pansorbin (Staph A) suspension was added to each tube, and after 15 minutes, the samples were centrifuged, washed twice, and the pellets counted. Nonspecific binding was determined by the addition of 130 nM unlabeled ZTNF4 to the ¹²⁵I-ZTNF4/TACI-Fc mix. Specific binding was calculated by subtracting the cpm bound in the presence of unlabeled ZTNF4 from the total cpm bound at each concentration of ¹²⁵I-ZTNF4. Each determination was performed in triplicate. Binding constants were calculated using GraphPad Prism software (Macintosh v. 3.0).

FIG. 4 illustrates the specific binding of ¹²⁵I-ZTNF4 to the various TACI-Fc constructs. Binding appeared to approach saturation with each construct, and was significantly higher for constructs TACI (d1-29)-Fc5, TACI (d1-29, d111-154)-Fc5, TACI (d1-29, d120-154)-Fc5, as compared with the binding of TACI (d1-29, d107-154)-Fc5. Bmax and Kd values were calculated, and the results are summarized in Table 7.

TABLE 7

Binding of ¹²⁵ I-ZTNF4 to TACI-Fc Constructs			
TACI-Fc Construct	Kd (nM)	Bmax (nM)	
TACI (d1-29)-Fc5 TACI (d1-29, d107-154)-Fc5	0.134 0.121	0.023 0.010	

TI IDEE / Continued				
Binding of ¹²⁵ I-ZTNF4 to TACI-Fc Constructs				
TACI-Fc Construct	Kd (nM)	Bmax (nM)		
TACI (d1-29, d111-154)-Fc5	0.115	0.018		
TACI (d1-29, d120-154)-Fc5	0.092	0.021		

Example 5

Measurement of Circulating ZTNF4

Levels of ZTNF4 in individuals with a disease condition (such as SLE, rheumatoid arthritis for example) relative to 15 normal individuals were determined using an electrochemiluminescence assay. A standard curve prepared from soluble, human ZTNF4 at 10 ng/ml, 1 ng/ml, 0.1 ng/ml, 0.01 ng/ml and 0 ng/ml was prepared in ORIGIN buffer (Igen, Gaithersburg, Md.). Serum samples were diluted in ORIGIN buffer. 20 The standards and samples were incubated at room temperature for two hours with biotinylated rabbit anti-human ZTNF4-NF BV antibody diluted to 1 μg/ml in Origin Assay Buffer (IGEN) and ruthenylated rabbit anti-human ZTNF4-NF BV polyclonal antibody diluted to 1 µg/ml in Origin 25 Assay Buffer (IGEN). Following the incubation the samples were vortexed and 0.4 mg/ml streptavidin Dynabeads (Dynal, Oslo, Norway) were added to each of the standards and samples at 50 µl/tube and incubated for 30 minutes at room temperature. Samples were then vortexed and samples were 30 read on an Origin Analyzer (Igen) according to manufacturer's instructions. The Origin assay is based on electrochemiluminescence and produces a readout in ECL. In one study, an elevated level of ZTNF4 was detected in the serum samples from both NZBWF1/J, and MRL/Mpj-Fas^{lpr} mice, which 35 have progressed to advanced stages of glomerulonephritis and autoimmune disease.

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The ORIGIN ASSAY was also used to measure levels of ZTNF4 in the blood of SLE patients, relative to circulating levels in normal individuals. A standard curve prepared from soluble, human ZTNF4 at 10 ng/ml, 1 ng/ml, 0.1 ng/ml, 0.01 ng/ml and 0 ng/ml was prepared in ORIGIN buffer (Igen). All patient samples were run in triplicate with a 25 µl final volume. The standards and samples were incubated at room temperature for two hours with a capture antibody, biotinylated rabbit anti-human ZTNF4-NF BV polyclonal antibody, diluted to 1 µg/ml in Origin Assay Buffer (IGEN) and a detection antibody, ruthenylated rabbit anti-human ZTNF4-NF BV polyclonal antibody, diluted to 1 µg/ml in Origin Assay Buffer (IGEN). Following the incubation the samples were vortexed, and 0.4 mg/ml streptavidin Dynabeads (Dynal) was added to each of the standards and samples at 50 μl/tube and incubated for 30 minutes at room temperature. Samples were then vortexed, and analyzed using an Origin 1.5 Analyzer (Igen) according to manufacturer's instructions.

This assay included 28 normal control samples and samples from 20 patients diagnosed with SLE. Elevated levels of ZTNF4 were observed in the serum of patients diagnosed with SLE, as compared with normal control serum donors. ZTNF4 levels were calculated as a fold increase of ZTNF4 levels in the patient or control samples as compared to an arbitrary human reference serum sample. The average of the 28 control samples was 1.36 fold over the human reference sample and the average of the 20 SLE patient samples was 4.92. Seven out of the 20 SLE patients had ZTNF4 levels that were two fold over the average of the control samples, whereas there was only one control individual that had a greater than two fold level over the control average.

From the foregoing, it will be appreciated that, although specific embodiments of the invention have been described herein for purposes of illustration, various modifications may be made without deviating from the spirit and scope of the invention. Accordingly, the invention is not limited except as by the appended claims.

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Gly	Val	Tyr	Leu	Leu 165	Tyr	Ser	Gln	Val	Leu 170	Phe	Gln	Asp	Val	Thr 175	Phe	
Thr	Met	Gly	Gln 180	Val	Val	Ser	Arg	Glu 185	Gly	Gln	Gly	Arg	Gln 190	Glu	Thr	
Leu	Phe	Arg 195	Сла	Ile	Arg	Ser	Met 200	Pro	Ser	His	Pro	Asp 205	Arg	Ala	Tyr	
Asn	Ser 210	Сув	Tyr	Ser	Ala	Gly 215	Val	Phe	His	Leu	His 220	Gln	Gly	Asp	Ile	
Leu 225	Ser	Val	Ile	Ile	Pro 230	Arg	Ala	Arg	Ala	Lys 235	Leu	Asn	Leu	Ser	Pro 240	
His	Gly	Thr	Phe	Leu 245	Gly	Phe	Val	Lys	Leu 250							
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ggat	icc a				ctg t Leu :											48
	tgg Trp															96
	ccg Pro															144
	ccc Pro															192
	aca Thr															240
	aac Asn 80															288
	cgg Arg			Gln		Asn			Tyr		Val					336
	gtc Val															384
	tcc Ser															432
	aaa Lys															480
	gat Asp 160															528
	ttc Phe															576
ccg	gag	aac	aac	tac	aag	acc	acg	cct	ccc	gtg	ctg	gac	tcc	gac	ggc	624

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P	ro (lu	Asn	Asn	Tyr 195	Lys	Thr	Thr	Pro	Pro 200	Val	Leu	Asp	Ser	Asp 205	Gly	
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										atg Met							720
	is :									tct Ser				tga			762
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				NCE :		-	-										
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Va	al I	Leu	Ser	Glu 20	Pro	Lys	Ser	Cys	Asp 25	Lys	Thr	His	Thr	Cys	Pro	Pro	
C	ys I	?ro	Ala 35	Pro	Glu	Leu	Leu	Gly 40	Gly	Pro	Ser	Val	Phe 45	Leu	Phe	Pro	
Pi		20 Jàa	Pro	ГÀа	Asp	Thr	Leu 55	Met	Ile	Ser	Arg	Thr 60	Pro	Glu	Val	Thr	
6! 6!		/al	Val	Val	Asp	Val 70	Ser	His	Glu	Asp	Pro 75	Glu	Val	Lys	Phe	Asn 80	
T	rp :	Гуr	Val	Asp	Gly 85	Val	Glu	Val	His	Asn 90	Ala	ГÀа	Thr	Lys	Pro 95	Arg	
				100					105	Val				110			
			115					120		Glu			125				
	-	L30					135			Lys		140					
14	15					150				Thr	155					160	
				-	165					Thr 170	=			-	175		
				180					185	Glu				190			
			195					200		Lys			205				
		210	-1-		-15		215		~P	-15		220	1			1	
	sn V 25	/al	Phe	Ser	Сув	Ser 230	Val	Met	His	Glu	Ala 235	Leu	His	Asn	His	Tyr 240	
Tì	ar (3ln	ГЛа	Ser	Leu 245	Ser	Leu	Ser	Pro	Gly 250	Lys						
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Phe Arg Arg
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agetgetett getgeatttg etetggaatt ettgtagaga tattaettgt eetteeagge	180
tgttctttct gtagctccct tgttttcttt ttgtgatc atg ttg cag atg gct ggg $$\operatorname{Met}$$ Leu Gln Met Ala Gly 1	236
cag tgc tcc caa aat gaa tat ttt gac agt ttg ttg cat gct tgc ata Gln Cys Ser Gln Asn Glu Tyr Phe Asp Ser Leu Leu His Ala Cys Ile 10 15 20	284
cct tgt caa ctt cga tgt tct tct aat act cct cct cta aca tgt cag Pro Cys Gln Leu Arg Cys Ser Ser Asn Thr Pro Pro Leu Thr Cys Gln 25 30 35	332
cgt tat tgt aat gca agt gtg acc aat tca gtg aaa gga acg aat gcg Arg Tyr Cys Asn Ala Ser Val Thr Asn Ser Val Lys Gly Thr Asn Ala 40 45 50	380
att ctc tgg acc tgt ttg gga ctg agc tta ata att tct ttg gca gtt Ile Leu Trp Thr Cys Leu Gly Leu Ser Leu Ile Ile Ser Leu Ala Val 55 60 65 70	428
ttc gtg cta atg ttt ttg cta agg aag ata agc tct gaa cca tta aag Phe Val Leu Met Phe Leu Leu Arg Lys Ile Ser Ser Glu Pro Leu Lys 75 80 85	476
gac gag ttt aaa aac aca gga tca ggt ctc ctg ggc atg gct aac att Asp Glu Phe Lys Asn Thr Gly Ser Gly Leu Leu Gly Met Ala Asn Ile 90 95 100	524
gac ctg gaa aag agc agg act ggt gat gaa att att ctt ccg aga ggc Asp Leu Glu Lys Ser Arg Thr Gly Asp Glu Ile Ile Leu Pro Arg Gly 105 110 115	572
ctc gag tac acg gtg gaa gaa tgc acc tgt gaa gac tgc atc aag agc Leu Glu Tyr Thr Val Glu Glu Cys Thr Cys Glu Asp Cys Ile Lys Ser 120 125 130	620
aaa ccg aag gtc gac tct gac cat tgc ttt cca ctc cca gct atg gag Lys Pro Lys Val Asp Ser Asp His Cys Phe Pro Leu Pro Ala Met Glu 135	668
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		35					40					45				
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Ile 65	Ile	Ser	Leu	Ala	Val 70	Phe	Val	Leu	Met	Phe 75	Leu	Leu	Arg	Lys	Ile 80	
Ser	Ser	Glu	Pro	Leu 85	Lys	Asp	Glu	Phe	Lys 90	Asn	Thr	Gly	Ser	Gly 95	Leu	
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Glu	Asp 130		Ile	ГÀа	Ser	Lys 135	Pro	Lys	Val	Asp	Ser 140	Asp	His	СЛа	Phe	
Pro 145	Leu	Pro	Ala	Met	Glu 150	Glu	Gly	Ala	Thr	Ile 155	Leu	Val	Thr	Thr	Lys 160	
Thr	Asn	Asp	Tyr	Сув 165	Lys	Ser	Leu	Pro	Ala 170	Ala	Leu	Ser	Ala	Thr 175	Glu	
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aga Arg 15 cca Pro	tgg Trp ccg Pro	Met I gtc Val tgc Cys	ctg Leu cca Pro	tcc Ser gca Ala	gag Glu 20 cct Pro	CCC Pro gaa Glu	aga Arg ctc Leu	tct Ser ctg Leu	tca Ser 999 Gly 40	gac Asp 25 gga Gly	Leu 10 aaa Lys ccg Pro	act Thr tca Ser	cac His gtc Val	aca Thr ttc Phe 45	tgc Cys 30 ctc Leu	96
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aga Arg 15 cca Pro ttc Phe gtc Val ttc Phe	tgg Trp ccc Pro aca Thr aac Asn 80	Met I I 1 gtc Val tgc Cys cca Pro tgc Cys 65 tgg Trp gag	ctg Leu cca Pro aaa Lys 50 gtg Val tac Tyr	tcc Ser gca Ala 35 ccc Pro gtg Val	gag Glu 20 cct Pro aag Lys gtg Val gac Asp	Trp I I 5 ccc Pro gaa Glu gac Asp gac Asp ggc Gly 85 aac	aga Arg ctc Leu acc Thr gtg Val 70 gtg Val agc	tct Ser ctg Leu ctc Leu 55 agc Ser gag Glu acg	tca Ser 999 Gly 40 atg Met cac His	gac Asp 25 gga Gly atc Ile gaa Glu cat His	Let 1 10 aaaa Lys ccg Pro tcc Ser gac Asp aat Asn 90 gtg	act Thr tca Ser cgg Arg cct Pro 75 gcc Ala	cac His gtc Val acc Thr 60 gag Glu aagg Lys	Ala I aca Thr ttc Phe 45 cct Pro gtc Val aca Thr	tgc Cys 30 ctc Leu gag Glu aag Lys	96 144 192 240
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aga Arg 15 cca Pro ttc Phe gtc Val ttc Phe ccg Pro 95 acc Thr	tgg Trp ccg Pro ccc Pro aca Thr acc Asn so cgg Arg gtc Val	Met I 1 1 gtc Val 1 tgc Cys cca Pro tgc Cys 65 tgg Trp gag Glu ctg Leu aac	ctg Leu cca Pro aaa Lys 50 gtg Val tac Tyr gag Glu cac His	tcc Ser gca Ala 35 ccc Pro gtg Val gtg Gln cag Gln	gag Glu 20 cct Pro aag Lys yal gac Asp tac Tyr 100 gac Asp ctc	Trp I I I Trp I I I I I I I I I I I I I I I I I I I	aga Arg ctc Leu acc Thr gtg Val 70 gtg Val agc Ser ctg Leu gcc	tct Ser ctg Leu ctc Leu 55 agc Ser gag Glu acg Thr aat Asn ccc	tca Ser ggg Gly 40 atg Met cac His gtg Val tac Tyr ggc Gly 120 atc	gac Asp 25 gga Gly atc Ile gaa Glu cat His Arg 105 aag Lys	Leu ville in the control of the cont	act Thr tca Ser cgg Arg cct Pro 75 gcc Ala gtc Val tac	cac His gtc Val acc Thr 60 gag Glu aag Lys agc ser	Ala I aca Thr ttc Phe 45 cct Pro gtc Val aca Thr gtc Val tgc Cys 125 tcc	tgc Cys 30 ctc Leu gag Glu aag Lys ctc Leu 110 aag	96 144 192 240 288

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	g gag o Glu															624
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Va:	l Leu	Ser	Glu 20	Pro	Arg	Ser	Ser	Asp 25	Lys	Thr	His	Thr	30 Cys	Pro	Pro	
Су	s Pro	Ala 35	Pro	Glu	Leu	Leu	Gly 40	Gly	Pro	Ser	Val	Phe 45	Leu	Phe	Pro	
Pro	50 50	Pro	Lys	Asp	Thr	Leu 55	Met	Ile	Ser	Arg	Thr 60	Pro	Glu	Val	Thr	
Су: 65	s Val	Val	Val	Asp	Val 70	Ser	His	Glu	Asp	Pro 75	Glu	Val	Lys	Phe	Asn 80	
Trj	o Tyr	Val	Asp	Gly 85	Val	Glu	Val	His	Asn 90	Ala	Lys	Thr	Lys	Pro 95	Arg	
Glı	u Glu	Gln	Tyr 100	Asn	Ser	Thr	Tyr	Arg 105	Val	Val	Ser	Val	Leu 110	Thr	Val	
Lei	u His	Gln 115	Asp	Trp	Leu	Asn	Gly 120	Lys	Glu	Tyr	ГÀа	Сув 125	ГÀа	Val	Ser	
Ası	n Lys 130		Leu	Pro	Ala	Pro 135	Ile	Glu	Lys	Thr	Ile 140	Ser	Lys	Ala	Lys	
Gl; 14!	y Gln 5	Pro	Arg	Glu	Pro 150	Gln	Val	Tyr	Thr	Leu 155	Pro	Pro	Ser	Arg	Asp 160	
Glı	u Leu	Thr	Lys	Asn 165	Gln	Val	Ser	Leu	Thr 170	Сув	Leu	Val	Lys	Gly 175	Phe	
Ту	r Pro	Ser	Asp 180	Ile	Ala	Val	Glu	Trp 185	Glu	Ser	Asn	Gly	Gln 190	Pro	Glu	
Ası	n Asn	Tyr 195	Lys	Thr	Thr	Pro	Pro 200	Val	Leu	Asp	Ser	Asp 205	Gly	Ser	Phe	
Ph	e Leu 210	_	Ser	Lys	Leu	Thr 215	Val	Asp	Lys	Ser	Arg 220	Trp	Gln	Gln	Gly	
As:	n Val	Phe	Ser	Cys	Ser 230	Val	Met	His	Glu	Ala 235	Leu	His	Asn	His	Tyr 240	

Thr Gln Lys Ser Leu Ser Leu Ser Pro Gly Lys

_														
				245				250						
<213 <213 <213 <220 <223 <223	1 > L1 2 > T 3 > O1 0 > F1 1 > N2 2 > L0	EATUI AME/I OCAT:	H: 70 DNA ISM: RE: KEY: ION:	62 Homo CDS (7)	(ed in	nmun	oglol	buli	n mo	iety		
< 400	0 > S1	EQUEI	NCE :	30										
ggat		-	_		_		ttc (Phe 1		_	_			-	48
							tct Ser							96
							gag Glu							144
							ctc Leu 55							192
							agc Ser							240
							gag Glu							288
							acg Thr							336
							aat Asn							384
							tcc Ser 135							432
							cag Gln							480
							gtc Val							528
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<212> TYPE: PRT
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Val Leu Ser Glu Pro Arg Ser Ser Asp Lys Thr His Thr Cys Pro Pro
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Cys Pro Ala Pro Glu Ala Glu Gly Ala Pro Ser Val Phe Leu Phe Pro
Pro Lys Pro Lys Asp Thr Leu Met Ile Ser Arg Thr Pro Glu Val Thr
Cys Val Val Val Asp Val Ser His Glu Asp Pro Glu Val Lys Phe Asn
Trp Tyr Val Asp Gly Val Glu Val His Asn Ala Lys Thr Lys Pro Arg
                      90
Glu Glu Gln Tyr Asn Ser Thr Tyr Arg Val Val Ser Val Leu Thr Val
         100
                           105
Leu His Gln Asp Trp Leu Asn Gly Lys Glu Tyr Lys Cys Lys Val Ser
                        120
Asn Lys Ala Leu Pro Ser Ser Ile Glu Lys Thr Ile Ser Lys Ala Lys
             135
Gly Gln Pro Arg Glu Pro Gln Val Tyr Thr Leu Pro Pro Ser Arg Asp
           150
                                  155
Glu Leu Thr Lys Asn Gln Val Ser Leu Thr Cys Leu Val Lys Gly Phe
              165
                                170
Tyr Pro Ser Asp Ile Ala Val Glu Trp Glu Ser Asn Gly Gln Pro Glu
Asn Asn Tyr Lys Thr Thr Pro Pro Val Leu Asp Ser Asp Gly Ser Phe
               200
Phe Leu Tyr Ser Lys Leu Thr Val Asp Lys Ser Arg Trp Gln Gln Gly
            215
Asn Val Phe Ser Cys Ser Val Met His Glu Ala Leu His Asn His Tyr
                230
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Thr Gln Lys Ser Leu Ser Leu Ser Pro Gly Lys
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<211> LENGTH: 762
<212> TYPE: DNA
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<222> LOCATION: (7)...(759)
<223> OTHER INFORMATION: Modified immunoglobulin moiety.
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aga tgg gtc ctg tcc gag ccc aaa tct tca gac aaa act cac aca tgc
                                                                 96
Arg Trp Val Leu Ser Glu Pro Lys Ser Ser Asp Lys Thr His Thr Cys
cca ccg tgc cca gca cct gaa gcc gag ggg gca ccg tca gtc ttc ctc
                                                                144
Pro Pro Cys Pro Ala Pro Glu Ala Glu Gly Ala Pro Ser Val Phe Leu
               35
                                 40
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Phe Pro Pro Lys Pro Lys Asp Thr Leu Met Ile Ser Arg Thr Pro Glu 50 55 60	192
gtc aca tgc gtg gtg gtg gac gtg agc cac gaa gac cct gag gtc aag Val Thr Cys Val Val Val Asp Val Ser His Glu Asp Pro Glu Val Lys 65 70 75	240
ttc aac tgg tac gtg gac ggc gtg gag gtg cat aat gcc aag aca aag Phe Asn Trp Tyr Val Asp Gly Val Glu Val His Asn Ala Lys Thr Lys 80 85 90	288
ccg cgg gag gag cag tac aac agc acg tac cgt gtg gtc agc gtc ctc Pro Arg Glu Glu Gln Tyr Asn Ser Thr Tyr Arg Val Val Ser Val Leu 95	336
acc gtc ctg cac cag gac tgg ctg aat ggc aag gag tac aag tgc aag Thr Val Leu His Gln Asp Trp Leu Asn Gly Lys Glu Tyr Lys Cys Lys 115 120 125	384
gtc tcc aac aaa gcc ctc cca tcc tcc atc gag aaa acc atc tcc aaa Val Ser Asn Lys Ala Leu Pro Ser Ser Ile Glu Lys Thr Ile Ser Lys 130 135 140	432
gcc aaa ggg cag ccc cga gaa cca cag gtg tac acc ctg ccc cca tcc Ala Lys Gly Gln Pro Arg Glu Pro Gln Val Tyr Thr Leu Pro Pro Ser 145 150 155	480
cgg gat gag ctg acc aag aac cag gtc agc ctg acc tgc ctg gtc aaa Arg Asp Glu Leu Thr Lys Asn Gln Val Ser Leu Thr Cys Leu Val Lys 160 165 170	528
ggc ttc tat ccc agc gac atc gcc gtg gag tgg gag agc aat ggg cag Gly Phe Tyr Pro Ser Asp Ile Ala Val Glu Trp Glu Ser Asn Gly Gln 175 180 185 190	576
ccg gag aac aac tac aag acc acg cct ccc gtg ctg gac tcc gac ggc Pro Glu Asn Asn Tyr Lys Thr Thr Pro Pro Val Leu Asp Ser Asp Gly 195 200 205	624
tcc ttc ttc ctc tac agc aag ctc acc gtg gac aag agc agg tgg cag Ser Phe Phe Leu Tyr Ser Lys Leu Thr Val Asp Lys Ser Arg Trp Gln 210 215 220	672
cag ggg aac gtc ttc tca tgc tcc gtg atg cat gag gct ctg cac aac Gln Gly Asn Val Phe Ser Cys Ser Val Met His Glu Ala Leu His Asn 225 230 235	720
cac tac acg cag aag agc ctc tcc ctg tct ccg ggt aaa tga His Tyr Thr Gln Lys Ser Leu Ser Leu Ser Pro Gly Lys 240 245 250	762
<210> SEQ ID NO 33 <211> LENGTH: 251 <212> TYPE: PRT <213> ORGANISM: Homo sapiens	
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Val Leu Ser Glu Pro Lys Ser Ser Asp Lys Thr His Thr Cys Pro Pro 20 25 30	
Cys Pro Ala Pro Glu Ala Glu Gly Ala Pro Ser Val Phe Leu Phe Pro 35 40 45	
Pro Lys Pro Lys Asp Thr Leu Met Ile Ser Arg Thr Pro Glu Val Thr 50 55 60	
Cys Val Val Val Asp Val Ser His Glu Asp Pro Glu Val Lys Phe Asn 65 70 75 80	
Trp Tyr Val Asp Gly Val Glu Val His Asn Ala Lys Thr Lys Pro Arg 85 90 95	

Glu Glu Gln Tyr Asn Ser Thr Tyr Arg Val Val Ser Val Leu Thr Val

			100					105					110				
Leu	His	Gln		Trp	Leu	Asn	Gly		Glu	Tyr	Lys	Cys		Val	Ser		
		115	•	-			120	1		•	•	125	1				
Asn	Lys 130	Ala	Leu	Pro	Ser	Ser 135	Ile	Glu	ГÀв	Thr	Ile 140	Ser	ГЛа	Ala	Lys		
	Gln	Pro	Arg	Glu	Pro	Gln	Val	Tyr	Thr		Pro	Pro	Ser	Arg	_		
145	T	m1	T	7	150	77-7	G	T	m1	155	T	77-7	T	G1	160 Db-		
GIU	ьeu	Tnr	гув	165	Gln	vai	ser	ьeu	170	Cys	ьeu	vai	гуз	175	Pne		
Tyr	Pro	Ser	Asp 180	Ile	Ala	Val	Glu	Trp 185	Glu	Ser	Asn	Gly	Gln 190	Pro	Glu		
Asn	Asn	Tyr 195	Lys	Thr	Thr	Pro	Pro 200	Val	Leu	Asp	Ser	Asp 205	Gly	Ser	Phe		
Phe	Leu 210	Tyr	Ser	ГÀа	Leu	Thr 215	Val	Asp	Lys	Ser	Arg 220	Trp	Gln	Gln	Gly		
Asn 225	Val	Phe	Ser	CAa	Ser 230	Val	Met	His	Glu	Ala 235	Leu	His	Asn	His	Tyr 240		
Thr	Gln	Lys	Ser	Leu 245	Ser	Leu	Ser	Pro	Gly 250	Lys							
)	יקווסק															
	ccc a	atg a	_	cac (ctg t Leu :					_							48
ggat aga	tgg	atg a Met 1 1 gtc	aag (Lys 1	cac (His D	_	Frp F 5 ccc	Phe I	Phe l	Leu I tca	Leu I gac	Leu V 10 aaa	Val i	Ala A	Ala E aca	Pro tgc		48 96
ggai aga Arg 15 cca	tgg Trp	atg a Met 1 1 gtc Val	aag (Lys) ctg Leu cca	cac (His l tcc Ser	Leu : gag Glu	Trp F 5 ccc Pro	Phe I aaa Lys gcc	tct Ser gag	tca Ser 999	gac Asp 25 gca	eu V 10 aaa Lys ccg	act Thr	cac His	aca Thr	tgc Cys 30	:	
aga Arg 15 cca Pro	tgg Trp ccg Pro	atg a Met I 1 gtc Val tgc Cys	ctg Lys l ctg Leu cca Pro	tcc Ser gca Ala 35	gag Glu 20	CCC Pro gaa Glu	aaa Lys gcc Ala	tct Ser gag Glu	tca Ser 999 Gly 40	gac Asp 25 gca Ala	Leu V 10 aaa Lys ccg Pro	act Thr tca Ser	cac His gtc Val	aca Thr ttc Phe 45	tgc Cys 30 ctc Leu		96
gga ¹ aga Arg 15 cca Pro ttc	tgg Trp ccg Pro	atg affect of the second secon	ctg Leu cca Pro aaa Lys 50	tcc Ser gca Ala 35 ccc Pro	gag Glu 20 cct Pro	CCC Pro gaa Glu gac Asp	aaa Lys gcc Ala acc Thr	tct Ser gag Glu ctc Leu 55	tca Ser 999 Gly 40 atg Met	gac Asp 25 gca Ala atc Ile	Leu V 10 aaa Lys ccg Pro tcc Ser	act Thr tca Ser cgg Arg	cac His gtc Val acc Thr 60	aca Thr ttc Phe 45 cct Pro	tgc Cys 30 ctc Leu gag Glu	:	96 144
aga Arg 15 cca Pro ttc Phe	tgg Trp ccg Pro ccc Thr	tgc Cys cca Pro	ctg Ctg Leu cca Pro aaaa Lys 50 gtg Val	tcc Ser gca Ala 35 ccc Pro gtg Val	gag Glu 20 cct Pro aag Lys	Trp I Ccc Pro gaa Glu gac Asp gac	aaa Lys gcc Ala acc Thr gtg Val 70 gtg	tct Ser gag Glu ctc Leu 55 agc Ser	tca Ser 999 Gly 40 atg Met cac His	gac Asp 25 gca Ala atc Ile gaa Glu cat	eu V 10 aaaa Lys ccg Pro tcc Ser gac Asp	act Thr tca Ser cgg Arg	cac His gtc Val acc Thr 60 gag Glu aag	Ala I aca Thr ttc Phe 45 cct Pro	tgc Cys 30 ctc Leu gag Glu aag Lys	2	96 144 192
ggai aga Arg 15 cca Pro ttc Phe gtc Val	tgg Trp ccg Pro ccc Thr	atg a Met 1 1 gtc Val tgc Cys cca Pro tgc Cys 65 tgg Trp gag	ctg Leu cca Pro aaaa Lys 50 gtg Val tac Tyr	tcc Ser gca Ala 35 ccc Pro gtg Val	gag Glu 20 cct Pro aag Lys gtg Val	Trp I 5 ccc Pro gaa Glu gac Asp gac Asp ggc Gly 85 aac	aaaa Lys gcc Ala acc Thr gtg Val 70 gtg Val	tct Ser gag Glu ctc Leu 55 agc Ser gag Glu acg	tca Ser ggg Gly 40 atg Met cac His	gac Asp 25 gca Ala atc Ile gaa Glu cat His	eu V 10 aaaa Lys ccg Pro tcc Ser gac Asp	act Thr tca Ser cgg Arg cct Pro 75 gcc Ala	cac His gtc Val acc Thr 60 gag Glu aagc Lys	Ala F aca Thr ttc Phe 45 cct Pro gtc Val aca Thr	tgc Cys 30 ctc Leu gag Glu aag Lys	:	96 144 192 240
aga Arg 15 cca Pro ttc Phe ttc Phe ccg Pro 95 acc	tgg Trp ccg Pro ccc Pro aca Thr acc Asn so cgg Arg	tgc Cys cca Pro tgc Cys 65 tgg Trp gag Glu ctg	ctg Leu cca Pro aaaa Lys 50 gtg Val tac Tyr gag Glu cac	tcc Ser gca Ala 35 ccc Pro gtg Val gtg Cal	gag Glu 20 cct Pro aag Lys gtg Val gac Asp	Trp I F 5 ccc Pro gaa Glu gac Asp gac Asp ggc Gly 85 aac Asn	aaaa Lys gcc Ala acc Thr gtg Val 70 gtg Val agc Ser	Phe I tott Ser gag Glu ctc Leu 55 agc Ser gag Glu acg Thr	tca Ser 999 Gly 40 atg Met Cac His 9tg Val	gac Asp 25 gca Ala atc Ile gaa Glu cat His cgt Arg 105 aag	Leu V 10 aaaa Lys ccg Pro tcc Ser gac Asp aat Asn 90 gtg Val	act Thr tca Ser cgg Arg cet Pro 75 gcc Ala	cac His gtc Val acc Thr 60 gag Glu aag Lys agc Ser aag	aca Thr ttc Phe 45 cct Pro gtc Val aca Thr	tgc Cys 30 ctc Leu gag Glu aag Lys ctc Leu	:	96 144 192 240
aga Arg 15 cca Pro ttc Phe gtc Val ttc Phe ccg Pro 95 acc Thr	tgg Trp ccg Pro ccc Pro aca Thr aac Asn 80 cgg Arg	tgc Cys cca Pro tgc Cys 65 tgg Trp gag Glu ctg Leu	ctg Leu cca Pro aaaa Lys 50 gtg Val tac Tyr gag Glu cac His	tcc Ser gca Ala 35 ccc Pro gtg Val gtg Gln cag Gln 115 gcc	gag Glu 20 cct Pro aag Lys gtg Val gac Asp tac Tyr 100 gac	Trp I F 5 ccc Pro gaa Glu gac Asp gac Asp ggc Gly 85 aac Asn tgg Trp cca	Phe I aaaa Lys gcc Ala acc Thr gtg Val 70 gtg Val agc ser ctg Leu	Phe I tot Ser gag Glu otc Leu 55 agc Ser gag Glu acg Thr aat Asn	tca Ser 999 Gly 40 atg Met Cac His gtg Val tac Tyr 990 Gly 120 atc	gac Asp 25 gca Ala atc Ile gaa Glu cat His Arg 105 aag Lys	Leu V 10 aaaa Lys ccg Pro tcc Ser gac Asp aat Asn 90 gtg Val gag Glu aaa	act Thr tca Ser cgg Arg Arg ctt Pro 75 gcc Ala gtc Val tac Tyr acc	cac His gtc Val acc Thr 60 gag Glu aag Lys agc ser	aca Thr ttc Phe 45 cct Pro gtc Val aca Thr gtc Val tgc Cys 125 tcc	tgc Cys 30 ctc Leu gag Glu aag Lys ctc Leu 110 aag Lys aaa	2	96 144 192 240 288

-continued

	- COI	ntinued
145	150 155	5
Arg Asp Glu Leu Thr Lys	aac cag gtc agc ctg acc tgo Asn Gln Val Ser Leu Thr Cys 165 170	
	atc gcc gtg gag tgg gag ago (le Ala Val Glu Trp Glu Sen 185	
	acc acg cct ccc gtg ctg gad Thr Thr Pro Pro Val Leu Asp 200	
_	aag ctc acc gtg gac aag ago Lys Leu Thr Val Asp Lys Ser 215	
	tgc tcc gtg atg cat gag gct Cys Ser Val Met His Glu Ala 230 235	a Leu His Asn
His Tyr Thr Gln Lys Ser	etc tcc ctg tct ccg ggt tga Leu Ser Leu Ser Pro Gly 245 250	a 759
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Cys Pro Ala Pro Glu Ala 35	Glu Gly Ala Pro Ser Val Pho 40 45	e Leu Phe Pro
	Leu Met Ile Ser Arg Thr Pro 55 60	o Glu Val Thr
Cys Val Val Val Asp Val	Ser His Glu Asp Pro Glu Val 75	l Lys Phe Asn 80
Trp Tyr Val Asp Gly Val 85	Glu Val His Asn Ala Lys Thi 90	r Lys Pro Arg 95
Glu Glu Gln Tyr Asn Ser 100	Thr Tyr Arg Val Val Ser Val 105	l Leu Thr Val 110
Leu His Gln Asp Trp Leu 115	Asn Gly Lys Glu Tyr Lys Cys 120 129	
_	Ser Ile Glu Lys Thr Ile Ser 135 140	r Lys Ala Lys
Gly Gln Pro Arg Glu Pro 145 150	Gln Val Tyr Thr Leu Pro Pro 155	o Ser Arg Asp 160
Glu Leu Thr Lys Asn Gln 165	al Ser Leu Thr Cys Leu Val 170	l Lys Gly Phe 175
Tyr Pro Ser Asp Ile Ala 180	/al Glu Trp Glu Ser Asn Gly 185	y Gln Pro Glu 190
Asn Asn Tyr Lys Thr Thr 195	Pro Pro Val Leu Asp Ser Asp 200 209	· · · ·
	Thr Val Asp Lys Ser Arg Trp 215 220	p Gln Gln Gly
Asn Val Phe Ser Cys Ser 225 230	al Met His Glu Ala Leu His 235	s Asn His Tyr 240

			00110	Inaca
Thr Gln Lys	Ser Leu Ser 245	Leu Ser Pro	Gly 250	
<220> FEATU <221> NAME, <222> LOCAT <223> OTHER	TH: 762 : DNA IISM: Homo sa JRE: /KEY: CDS FION: (7)(R INFORMATION	759)	mmunoglobulin moie	ety.
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Met 1	Lys His Leu	Trp Phe Phe I 5	seu Leu Leu Val Al 10	a Ala Pro
		Pro Lys Ser	tgc gac aaa act c Cys Asp Lys Thr F 25	
			ggg gga ccg tca g Gly Gly Pro Ser V 40	
	-		atg atc tcc cgg a Met Ile Ser Arg T	5 5
	Val Val Val		cac gaa gac cct g His Glu Asp Pro C 75	
			gtg cat aat gcc a Val His Asn Ala I 90	
		Gln Ser Thr	tac cgt gtg gtc a Tyr Arg Val Val S 105	
			ggc aag gag tac a Gly Lys Glu Tyr I 120	
			atc gag aaa acc a Ile Glu Lys Thr 1	
	/ Gln Pro Arg		gtg tac acc ctg c Val Tyr Thr Leu F 155	
			agc ctg acc tgc c Ser Leu Thr Cys I 170	
		Ile Ala Val	gag tgg gag agc a Glu Trp Glu Ser A 185	
			ccc gtg ctg gac t Pro Val Leu Asp S 200	
	_	-	gtg gac aag agc a Val Asp Lys Ser A	
	n Val Phe Ser		atg cat gag gct o Met His Glu Ala I 235	=
		_	tct ccg ggt aaa t Ser Pro Gly Lys 250	ga 762

<210> SEQ ID NO 37 <211> LENGTH: 251 <212> TYPE: PRT <213 > ORGANISM: Homo sapiens <400> SEQUENCE: 37 Met Lys His Leu Trp Phe Phe Leu Leu Leu Val Ala Ala Pro Arg Trp Val Leu Ser Glu Pro Lys Ser Cys Asp Lys Thr His Thr Cys Pro Pro Cys Pro Ala Pro Glu Leu Leu Gly Gly Pro Ser Val Phe Leu Phe Pro Pro Lys Pro Lys Asp Thr Leu Met Ile Ser Arg Thr Pro Glu Val Thr Cys Val Val Val Asp Val Ser His Glu Asp Pro Glu Val Lys Phe Asn 65 70 75 80 Trp Tyr Val Asp Gly Val Glu Val His Asn Ala Lys Thr Lys Pro Arg Glu Glu Gln Tyr Gln Ser Thr Tyr Arg Val Val Ser Val Leu Thr Val 105 Leu His Gln Asp Trp Leu Asn Gly Lys Glu Tyr Lys Cys Lys Val Ser 120 Asn Lys Ala Leu Pro Ala Pro Ile Glu Lys Thr Ile Ser Lys Ala Lys Gly Gln Pro Arg Glu Pro Gln Val Tyr Thr Leu Pro Pro Ser Arg Asp 150 155 Glu Leu Thr Lys Asn Gln Val Ser Leu Thr Cys Leu Val Lys Gly Phe 170 Tyr Pro Ser Asp Ile Ala Val Glu Trp Glu Ser Asn Gly Gln Pro Glu 180 185 Asn Asn Tyr Lys Thr Thr Pro Pro Val Leu Asp Ser Asp Gly Ser Phe 200 Phe Leu Tyr Ser Lys Leu Thr Val Asp Lys Ser Arg Trp Gln Gln Gly 215 Asn Val Phe Ser Cys Ser Val Met His Glu Ala Leu His Asn His Tyr 230 Thr Gln Lys Ser Leu Ser Leu Ser Pro Gly Lys 245 <210> SEQ ID NO 38 <211> LENGTH: 762 <212> TYPE: DNA <213 > ORGANISM: Homo sapiens <220> FEATURE: <221> NAME/KEY: CDS <222> LOCATION: (7)...(759) <223> OTHER INFORMATION: Modified immunoglobulin moiety. <400> SEQUENCE: 38 Met Lys His Leu Trp Phe Phe Leu Leu Leu Val Ala Ala Pro 1.0 aga tgg gtc ctg tcc gag ccc aaa tct tca gac aaa act cac aca tgc 96 Arg Trp Val Leu Ser Glu Pro Lys Ser Ser Asp Lys Thr His Thr Cys 20 25 cca ccg tgc cca gca cct gaa ctc ctg ggg gga ccg tca gtc ttc ctc 144 Pro Pro Cys Pro Ala Pro Glu Leu Leu Gly Gly Pro Ser Val Phe Leu 40

						gac Asp										192
						gac Asp										240
						ggc Gly 85										288
						aac Asn										336
						tgg Trp										384
						cca Pro										432
						gaa Glu										480
						aac Asn 165										528
						atc Ile										576
_					_	acc Thr	_				_	_		_		624
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		_	_	_	_	ctc Leu 245		_		_			tga			762
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Val	Leu	Ser	Glu 20	Pro	Lys	Ser	Ser	Asp 25	Lys	Thr	His	Thr	30 Cys	Pro	Pro	
Cys	Pro	Ala 35	Pro	Glu	Leu	Leu	Gly 40	Gly	Pro	Ser	Val	Phe 45	Leu	Phe	Pro	
Pro	Lys 50	Pro	Lys	Asp	Thr	Leu 55	Met	Ile	Ser	Arg	Thr 60	Pro	Glu	Val	Thr	
Сув 65	Val	Val	Val	Asp	Val 70	Ser	His	Glu	Asp	Pro 75	Glu	Val	Lys	Phe	Asn 80	
Trp	Tyr	Val	Asp	Gly 85	Val	Glu	Val	His	Asn 90	Ala	Lys	Thr	Lys	Pro 95	Arg	

Glu Glu Gln Tyr Asn Ser Thr Tyr Arg Val Val Ser Val Leu Thr Val	
Leu His Gln Asp Trp Leu Asn Gly Lys Glu Tyr Lys Cys Lys Val Ser 115 120 125	
Asn Lys Ala Leu Pro Ala Pro Ile Glu Lys Thr Ile Ser Lys Ala Lys 130 135 140	
Gly Gln Pro Arg Glu Pro Gln Val Tyr Thr Leu Pro Pro Ser Arg Asp 145 150 155 160	
Glu Leu Thr Lys Asn Gln Val Ser Leu Thr Cys Leu Val Lys Gly Phe 165 170 175	
Tyr Pro Ser Asp Ile Ala Val Glu Trp Glu Ser Asn Gly Gln Pro Glu 180 185 190	
Asn Asn Tyr Lys Thr Thr Pro Pro Val Leu Asp Ser Asp Gly Ser Phe	
Phe Leu Tyr Ser Lys Leu Thr Val Asp Lys Ser Arg Trp Gln Gln Gly	
210 215 220 Asn Val Phe Ser Cys Ser Val Met His Glu Ala Leu His Asn His Tyr	
225 230 235 240 Thr Gln Lys Ser Leu Ser Pro Gly Lys	
245 250	
<210> SEQ ID NO 40 <211> LENGTH: 29	
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<210> SEQ ID NO 41	
<211> LENGTH: 29 <212> TYPE: DNA	
<213> ORGANISM: Homo sapiens	
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<211> LENGTH: 29 <212> TYPE: DNA	
<213> ORGANISM: Homo sapiens	
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<211> LENGTH: 31
<212> TYPE: DNA
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<220> FEATURE:
<223 > OTHER INFORMATION: PCR primer.
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<210> SEQ ID NO 45
<211> LENGTH: 30
<212> TYPE: DNA
<213 > ORGANISM: Homo sapiens
<220> FEATURE:
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<211> LENGTH: 28
<212> TYPE: DNA
<213 > ORGANISM: Homo sapiens
<220> FEATURE:
<223> OTHER INFORMATION: PCR primer.
<400> SEQUENCE: 46
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<210> SEQ ID NO 47
<211> LENGTH: 26
<212> TYPE: DNA
<213 > ORGANISM: Homo sapiens
<220> FEATURE:
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<400> SEQUENCE: 47
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<211> LENGTH: 28
<212> TYPE: DNA
<213> ORGANISM: Homo sapiens
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tgc cgc aag gag caa ggc aag ttc tat gac cat ctc ctg agg gac tgc Cys Arg Lys Glu Gln Gly Lys Phe Tyr Asp His Leu Leu Arg Asp Cys

atc agc tgt gcc tcc atc tgt gga cag cac cct aag caa tgt gca tac Ile Ser Cys Ala Ser Ile Cys Gly Gln His Pro Lys Gln Cys Ala Tyr

ttc tgt gag aac aag ctc agg agc cca gtg aac ctt cca cca gag ctc Phe Cys Glu Asn Lys Leu Arg Ser Pro Val Asn Leu Pro Pro Glu Leu

agg aga cag cag agt gga gaa gtt gaa aac aat tca gac aac tcg gga Arg Arg Gln Arg Ser Gly Glu Val Glu Asn Asn Ser Asp Asn Ser Gly 436 135 130

agg tac caa gga ttg gag cac aga ggc tca gaa gca agt cca gct ctc Arg Tyr Gln Gly Leu Glu His Arg Gly Ser Glu Ala Ser Pro Ala Leu 484 150

cca ggt ctc aag gag ccc aaa tct tca gac aaa act cac aca tgc cca Pro Gly Leu Lys Glu Pro Lys Ser Ser Asp Lys Thr His Thr Cys Pro 532 160 165

ccg tgc cca gca cct gaa gcc gag ggg gca ccg tca gtc ttc ctc ttc 580 Pro Cys Pro Ala Pro Glu Ala Glu Gly Ala Pro Ser Val Phe Leu Phe 180

ccc cca aaa ccc aag gac acc ctc atg atc tcc cgg acc cct gag gtc 628 Pro Pro Lys Pro Lys Asp Thr Leu Met Ile Ser Arg Thr Pro Glu Val 190 195 200

676

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772 cgg gag gag cag tac aac agc acg tac cgt gtg gtc agc gtc ctc acc Arg Glu Glu Gln Tyr Asn Ser Thr Tyr Arg Val Val Ser Val Leu Thr

gtc ctg cac cag gac tgg ctg aat ggc aag gag tac aag tgc aag gtc Val Leu His Gln Asp Trp Leu Asn Gly Lys Glu Tyr Lys Cys Lys Val

tcc aac aaa gcc ctc cca tcc tcc atc gag aaa acc atc tcc aaa gcc 868 Ser Asn Lys Ala Leu Pro Ser Ser Ile Glu Lys Thr Ile Ser Lys Ala

aaa ggg cag ccc cga gaa cca cag gtg tac acc ctg ccc cca tcc cgg 916 Lys Gly Gln Pro Arg Glu Pro Gln Val Tyr Thr Leu Pro Pro Ser Arg

295

gat gag ctg acc aag aac cag gtc agc ctg acc tgc ctg gtc aaa ggc Asp Glu Leu Thr Lys Asn Gln Val Ser Leu Thr Cys Leu Val Lys Gly 310

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290

3.05

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Glu	Pro	Lys	Ser	Ser 165	Asp	Lys	Thr	His	Thr 170	Сув	Pro	Pro	Сув	Pro 175	Ala	
Pro	Glu	Ala	Glu 180	Gly	Ala	Pro	Ser	Val 185	Phe	Leu	Phe	Pro	Pro 190	Lys	Pro	
Lys	Asp	Thr 195	Leu	Met	Ile	Ser	Arg 200	Thr	Pro	Glu	Val	Thr 205	Cha	Val	Val	
Val	Asp 210	Val	Ser	His	Glu	Asp 215	Pro	Glu	Val	Lys	Phe 220	Asn	Trp	Tyr	Val	
Asp 225	Gly	Val	Glu	Val	His 230	Asn	Ala	Lys	Thr	Lys 235	Pro	Arg	Glu	Glu	Gln 240	
Tyr	Asn	Ser	Thr	Tyr 245	Arg	Val	Val	Ser	Val 250	Leu	Thr	Val	Leu	His 255	Gln	
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Leu	Pro	Ser 275	Ser	Ile	Glu	Lys	Thr 280	Ile	Ser	Lys	Ala	Lys 285	Gly	Gln	Pro	
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		ctg Leu 210								676	
		aac Asn								724	
		Gly 999								772	
		gag Glu								820	
		tat Tyr								868	
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1082

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Gln	Gly	Lys	Phe	Tyr 85	Asp	His	Leu	Leu	Arg 90	Asp	CÀa	Ile	Ser	Сув 95	Ala	
Ser	Ile	Сув	Gly 100	Gln	His	Pro	Lys	Gln 105	Cys	Ala	Tyr	Phe	Cys 110	Glu	Asn	
Lys	Leu	Arg 115	Ser	Glu	Pro	Lys	Ser 120	Ser	Asp	Lys	Thr	His 125	Thr	Cys	Pro	
Pro	Cys 130		Ala	Pro	Glu	Ala 135	Glu	Gly	Ala	Pro	Ser 140	Val	Phe	Leu	Phe	
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Lys	Gly	Gln	Pro	Arg 245	Glu	Pro	Gln	Val	Tyr 250	Thr	Leu	Pro	Pro	Ser 255	Arg	
Asp	Glu	Leu	Thr 260		Asn	Gln	Val	Ser 265	Leu	Thr	Cys	Leu	Val 270	Lys	Gly	
Phe	Tyr	Pro 275	Ser	Asp	Ile	Ala	Val 280	Glu	Trp	Glu	Ser	Asn 285	Gly	Gln	Pro	
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					cgt Arg											148

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	aag Lys															532
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	agc Ser															1012
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Gln Arg Thr Cys Ala Ala Phe Cys Arg Ser Leu Ser Cys Arg Lys Glu
Gln Gly Lys Phe Tyr Asp His Leu Leu Arg Asp Cys Ile Ser Cys Ala
Ser Ile Cys Gly Gln His Pro Lys Gln Cys Ala Tyr Phe Cys Glu Asn
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Lys Leu Arg Ser Pro Val Asn Leu Pro Pro Glu Leu Arg Glu Pro Lys
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Glu Gly Ala Pro Ser Val Phe Leu Phe Pro Pro Lys Pro Lys Asp Thr
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Ser His Glu Asp Pro Glu Val Lys Phe Asn Trp Tyr Val Asp Gly Val
Glu Val His Asn Ala Lys Thr Lys Pro Arg Glu Glu Gln Tyr Asn Ser
Thr Tyr Arg Val Val Ser Val Leu Thr Val Leu His Gln Asp Trp Leu
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Gln Val Tyr Thr Leu Pro Pro Ser Arg Asp Glu Leu Thr Lys Asn Gln
Val Ser Leu Thr Cys Leu Val Lys Gly Phe Tyr Pro Ser Asp Ile Ala
Val Glu Trp Glu Ser Asn Gly Gln Pro Glu Asn Asn Tyr Lys Thr Thr
Pro Pro Val Leu Asp Ser Asp Gly Ser Phe Phe Leu Tyr Ser Lys Leu
        310
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Thr Val Asp Lys Ser Arg Trp Gln Gln Gly Asn Val Phe Ser Cys Ser
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gaacagatgg tecceagate ggteeegeee teageagttt etagagaace ateagatgtt	720
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60

That which is claimed:

- 1. A method of treating glomerulonephritis associated with 40 IgA nephropathy comprising administering to a mammalian subject in need thereof a composition comprising a transmembrane activator and calcium modulator and cyclophilin ligand-interactor (TACI)—immunoglobulin fusion protein, wherein the TACI-immunoglobulin fusion protein comprises:
 - (a) a TACI receptor moiety that consists of:
 - i) amino acid residues 30-110 of SEQ ID NO:2; or,
 - ii) amino acid residues 30-154 of SEQ ID NO:2;
 - wherein the TACI receptor moiety binds at least one of ZTNF2 or ZTNF4, and
 - (b) an immunoglobulin moiety comprising a C_{H2} and C_{H3} domain.
- 2. The method of claim 1, wherein the immunoglobulin $_{55}$ moiety is an IgG1 immunoglobulin moiety.
- 3. The method of claim 2, wherein the immunoglobulin moiety is an IgG1 Fc fragment that comprises a disulfide linked heavy chain hinge region, a C_{H2} domain and a C_{H3} domain.
- **4.** A method of treating glomerulonephritis associated with IgA nephropathy comprising administering to a mammalian subject in need thereof a composition comprising a transmembrane activator and calcium modulator and cyclophilin ligand-interactor (TACI)—immunoglobulin fusion protein, 65 wherein the TACI-immunoglobulin fusion protein comprises:

- (a) a TACI receptor moiety that consists of:
 - i) amino acid residues 30-110 of SEQ ID NO:2; or,
- ii) amino acid residues 30-154 of SEQ ID NO:2;
- wherein the TACI receptor moiety binds at least one of ZTNF2 or ZTNF4, and
- (b) an immunoglobulin moiety comprising a C_{H2} and C_{H3} domain,
- wherein the TACI-immunoglobulin fusion protein has an amino acid sequence of SEQ ID NO:54.
- 5. A method of treating glomerulonephritis associated with IgA nephropathy comprising administering to a mammalian subject in need thereof a composition comprising a transmembrane activator and calcium modulator and cyclophilin ligand-interactor (TACI)—immunoglobulin fusion protein, wherein the TACI-immunoglobulin fusion protein comprises:
 - (a) a TACI receptor moiety that consists of:
 - i) amino acid residues 30-110 of SEQ ID NO:2; or,
 - ii) amino acid residues 30-154 of SEQ ID NO:2;
 - wherein the TACI receptor moiety binds at least one of ZTNF2 or ZTNF4, and
 - (b) an immunoglobulin moiety comprising a C_{H2} and C_{H3} domain,
 - wherein the TACI-immunoglobulin fusion protein has an amino acid sequence comprising the secreted form of the amino acid sequence of SEQ ID NO:54.
- **6**. The method of claim **5**, wherein the TACI-immunoglobulin fusion protein comprises the amino acid sequence of SEQ ID NO:54, wherein the optimized tPA (otPA) leader sequence (SEQ ID NO:25) has been removed.

7. The method of claim 1, wherein the TACI-immunoglobulin fusion protein is a dimer.

8. The method of claim 1, wherein said subject is a human.

9. The method of claim 4, wherein said subject is a human.

10. The method of claim 5, wherein said subject is a human.

* * * * *